

UNPUBLISHED PRELIMINARY DATA

NATIONAL BUREAU OF STANDARDS REPORT

8595

PRELIMINARY REPORT ON A SURVEY OF
THERMODYNAMIC PROPERTIES OF THE COMPOUNDS OF THE ELEMENTS CHNOPS

N65 13448

FACILITY FORM 602

(ACCESSION NUMBER)

(THRU)

34

/

(PAGES)

(CODE)

CR-59922

33

(NASA CR OR TMX OR AD NUMBER)

(CATEGORY)

Progress Report for the Period 1 August to 31 October 1964

to

National Aeronautics and Space Administration

GPO PRICE \$ _____

OTS PRICE(S) \$ _____

Hard copy (HC) .00Microfiche (MF) .50

1 November 1964

U.S. DEPARTMENT OF COMMERCE
NATIONAL BUREAU OF STANDARDS

NATIONAL BUREAU OF STANDARDS REPORT

NBS PROJECT

221-11-0429

1 November 1964

NBS REPORT

8595

A SURVEY OF THERMODYNAMIC PROPERTIES OF THE COMPOUNDS OF THE ELEMENTS CHNOPS

George T. Armstrong, Eugene S. Domalski

George T. Furukawa and Mary A. Krivanec

Heat Division

Progress Report for the Period 1 August to 31 October 1964

to

NATIONAL AERONAUTICS AND SPACE ADMINISTRATION

Contract No. R-138

IMPORTANT NOTICE

NATIONAL BUREAU OF STANDARDS REPORTS are usually preliminary or progress accounting documents intended for use within the Government. Before material in the reports is formally published it is subjected to additional evaluation and review. For this reason, the publication, reprinting, reproduction, or open-literature listing of this Report, either in whole or in part, is not authorized unless permission is obtained in writing from the Office of the Director, National Bureau of Standards, Washington, D.C. 20234. Such permission is not needed, however, by the Government agency for which the Report has been specifically prepared if that agency wishes to reproduce additional copies for its own use.



U.S. DEPARTMENT OF COMMERCE

NATIONAL BUREAU OF STANDARDS

TABLE OF CONTENTS

	<u>Page</u>
Foreword	1
Introduction	1
Section I. Analysis of Heat-Capacity Data on Some Amino Acids	2
Section II. Heats and Free Energies of Formation of Compounds of C, H, N, O, P, and S	20

FOREWORD

A study at the National Bureau of Standards (NBS), of which this is the second progress report, has been undertaken to meet the need of the National Aeronautics and Space Administration (NASA) for thermodynamic information on biologically related materials important to the space program for several reasons. Among these reasons are the necessity of inferring the maximum amount of useful chemistry of incompletely accessible environments, for which only limited information is available, the possibility of the occurrence of organic compounds naturally synthesized under primitive conditions, and the possibility of theoretically recovering part of the prebiological history of the earth.

This program is being carried out under the technical supervision of Dr. George Jacobs of NASA, and with the consultation of Dr. Harold Morowitz of the Yale University, Department of Molecular Biology and Biophysics, and Dr. C. W. Beckett of the Heat Division (NBS). The contract (Contract No. R-138) was initiated May 1, 1964, and this report covers the second quarter of the term of the contract.

George T. Armstrong

George T. Armstrong
Supervisory Chemist
Project Leader

INTRODUCTION

Summary of Progress on the Survey of Thermodynamic Data

Following the plan outlined in the previous quarterly report, the survey of thermodynamic data on compounds of the elements C, H, N, O, P, and S was continued by two groups working in parallel. The work of one group is related to the heat capacities, enthalpies, entropies and free energies of the compounds of interest, while the work of the other group is concerned with enthalpy changes and free energy changes in reactions of the same classes of compounds.

The work of the first group was primarily devoted to an evaluation and smoothing of the available data on selected compounds for which thermodynamic data covering a range of temperatures had previously been found. The results of this evaluation is given in Section I of this report as a series of seventeen tables.

The work of the second group was devoted to an assessment of the data on the preliminary list of compounds containing one carbon atom or fewer per molecule, which is found in the appendix of the first quarterly report of this contract (NBS Report 8521). The search for this data was not entirely complete, but provides a coverage of the available data selected from several sources which perform an evaluation of existing data. The data found are shown in tabular form in Section II of this report as Table 1, preceded by a brief discussion of the completeness, and followed by a list of sources of the data.

Our consultant on this project, Dr. Harold Morowitz, agreed to provide a list of more complex organic compounds, ubiquitous to living organisms. (Reference, Memorandum of August 11, 1964, Appendix, NBS Report 8521) The preparation of this list is being handled as a purchase order. When received, the list will provide a basis for search for data of more complex organic compounds.

Section I

Analysis of Heat-Capacity Data on Some Amino Acids

George T. Furukawa and Mary A. Krivanec

The heat-capacity data on seventeen amino acids were examined and analyzed on the IBM 7094 computer. (Some of the data were on the hydrates and hydrachlorides of the amino acids.) Thermodynamic properties were calculated from the smoothed values of heat capacity and tables prepared from 0 to 300°K.

The molal values given are based on the 1961 atomic weights and the energy unit calorie is equal to 4.1840 joules. The references used in the data analysis are listed at the bottom of each table.

TABLE 1

MOLAL THERMODYNAMIC FUNCTIONS FOR GLYCINE
 $(\text{NH}_2\text{CH}_2\text{COOH})$
 SOLID PHASE

GRAM MOLECULAR WT. = 75.06765 GRAMS

T DEG K = 273.15 + T DEG C

1 CAL = 4.1840 JOULES

T DEG K	C _P CAL/DEG	(H _T ⁰ -H ₀ ^C) CAL	(H _T ⁰ -H ₀ ^C)/T CAL/DEG	S _T ⁰ CAL/DEG	-(α _T ⁰ -H ₀ ^C) CAL	-(α _T ⁰ -H ₀ ^C)/T CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.007	0.009	0.002	0.002	0.003	0.001
10.00	0.064	0.155	0.015	0.021	0.050	0.005
15.00	0.232	0.832	0.055	0.073	0.266	0.018
20.00	0.570	2.765	0.138	0.182	0.876	0.044
25.00	1.067	6.803	0.272	0.360	2.203	0.088
30.00	1.682	13.634	0.454	0.608	4.594	0.153
35.00	2.377	23.757	0.679	0.918	8.384	0.240
40.00	3.108	37.455	0.936	1.283	13.868	0.347
45.00	3.855	54.863	1.219	1.692	21.290	0.473
50.00	4.600	76.010	1.520	2.137	30.850	0.617
55.00	5.317	100.81	1.833	2.609	42.707	0.776
60.00	6.010	129.14	2.152	3.102	56.978	0.950
65.00	6.671	160.86	2.475	3.609	73.752	1.135
70.00	7.284	195.77	2.797	4.127	93.089	1.330
75.00	7.862	233.65	3.115	4.649	115.03	1.534
80.00	8.421	274.36	3.430	5.174	139.58	1.745
85.00	8.957	317.82	3.759	5.701	166.77	1.962
90.00	9.450	363.85	4.043	6.227	196.59	2.184
95.00	9.903	412.20	4.339	6.750	229.04	2.411
100.00	10.330	462.64	4.626	7.269	264.09	2.641
105.00	10.752	515.55	4.910	7.784	301.72	2.874
110.00	11.170	570.36	5.185	8.293	341.92	3.108
115.00	11.565	627.20	5.454	8.799	384.65	3.345
120.00	11.943	685.98	5.716	9.299	429.90	3.582
125.00	12.315	746.62	5.973	9.794	477.63	3.821
130.00	12.686	805.13	6.224	10.284	527.83	4.060
135.00	13.051	873.47	6.470	10.770	580.47	4.300
140.00	13.402	937.61	6.712	11.251	635.52	4.539
145.00	13.742	1007.9	6.948	11.727	692.97	4.779
150.00	14.078	1077.0	7.180	12.199	752.79	5.019
155.00	14.411	1146.2	7.408	12.666	814.95	5.258
160.00	14.737	1221.1	7.632	13.128	875.44	5.496
165.00	15.061	1295.6	7.852	13.587	946.23	5.735
170.00	15.387	1371.7	8.069	14.041	1015.3	5.972
175.00	15.710	1447.5	8.283	14.492	1086.6	6.209
180.00	16.027	1526.8	8.493	14.939	1160.2	6.446
185.00	16.338	1609.7	8.701	15.382	1236.0	6.681
190.00	16.650	1692.2	8.906	15.822	1314.0	6.916
195.00	16.965	1776.2	9.109	16.259	1394.2	7.150
200.00	17.281	1861.9	9.309	16.692	1476.6	7.383
205.00	17.592	1947.0	9.508	17.123	1561.2	7.615
210.00	17.898	2037.6	9.704	17.551	1647.8	7.847
215.00	18.201	2128.0	9.898	17.975	1736.7	8.077
220.00	18.505	2219.8	10.090	18.397	1827.6	8.307
225.00	18.818	2313.1	10.280	18.816	1920.6	8.536
230.00	19.133	2407.9	10.469	19.233	2015.7	8.764
235.00	19.447	2504.4	10.657	19.648	2113.0	8.991
240.00	19.768	2602.5	10.844	20.062	2212.2	9.218
245.00	20.072	2702.3	11.030	20.473	2313.6	9.443
250.00	20.458	2803.7	11.215	20.883	2417.0	9.668
255.00	20.795	2906.9	11.400	21.291	2522.4	9.892
260.00	21.132	3011.7	11.583	21.698	2629.9	10.115
265.00	21.472	3118.4	11.767	22.104	2739.4	10.337
270.00	21.811	3220.4	11.950	22.509	2850.9	10.555
273.15	22.023	3295.5	12.065	22.763	2922.2	10.698
275.00	22.148	3335.3	12.132	22.912	2964.5	10.780
280.00	22.482	3447.9	12.314	23.314	3080.0	11.000
285.00	22.814	3561.1	12.495	23.715	3197.6	11.220
290.00	23.144	3676.0	12.676	24.114	3317.2	11.439
295.00	23.475	3792.6	12.856	24.513	3438.7	11.657
300.00	23.687	3866.9	12.969	24.763	3516.3	11.794
301.15	23.814	3910.8	13.036	24.910	3562.3	11.874

^C IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Parks, G. S., Huffman, H. M. and Barnmore, M.,
 Thermal data on organic compounds. XI. The heat capacities, entropies
 and free energies of ten compounds containing oxygen or nitrogen,
J. Am. Chem. Soc., **55**, 2733-2740 (1933).

Hutchens, J. O., Cole, A. G. and Stout, J. W.,
 Heat capacities from 11 to 305°K. and entropies of L-alanine and glycine,
J. Am. Chem. Soc., **82**, 4813-4815 (1960).

TABLE 2

MOLAL THERMODYNAMIC FUNCTIONS FOR L-ALANINE
 $(\text{CH}_3(\text{NH}_2)\text{CHCOOH})$

SOLID PHASE

GRAM MOLECULAR WT. = 89.09474 GRAMS
 $T \text{ DEG K} = 273.15 + T \text{ DEG C}$

1 CAL = 4.1840 JOULES

T DEG K	c_p^C CAL/DEG	$(H_{T-H_0^C})$ CAL	$(H_{T-H_0^C})/T$ CAL/DEG	S_T^0 CAL/DEG	$-(\alpha_{T-H_0^C})$ CAL	$-(\alpha_{T-H_0^C})/T$ CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.015	0.018	0.004	0.005	0.006	0.001
10.00	0.117	0.293	0.029	0.039	0.098	0.010
15.00	0.401	1.494	0.100	0.133	0.496	0.033
20.00	0.919	4.701	0.235	0.314	1.571	0.079
25.00	1.610	10.970	0.439	0.590	3.793	0.152
30.00	2.414	20.991	0.700	0.954	7.620	0.254
35.00	3.289	35.233	1.007	1.391	13.453	0.384
40.00	4.153	53.853	1.346	1.687	21.628	0.541
45.00	5.004	76.745	1.705	2.425	32.393	0.720
50.00	5.840	103.87	2.077	2.496	45.935	0.919
55.00	6.639	135.08	2.456	3.590	62.393	1.134
60.00	7.412	170.21	2.837	4.201	81.866	1.364
65.00	8.155	209.15	3.218	4.824	104.43	1.607
70.00	8.838	251.65	3.595	5.454	130.12	1.859
75.00	9.503	297.51	3.967	6.086	158.97	2.120
80.00	10.186	346.73	4.334	6.721	190.99	2.387
85.00	10.850	399.33	4.698	7.359	226.19	2.661
90.00	11.450	455.11	5.057	7.996	264.58	2.940
95.00	12.005	513.76	5.408	8.631	306.15	3.223
100.00	12.552	575.15	5.752	9.260	350.88	3.509
105.00	13.093	639.27	6.088	9.886	398.74	3.798
110.00	13.612	706.04	6.419	10.507	449.73	4.088
115.00	14.112	775.36	6.742	11.123	503.80	4.381
120.00	14.603	847.15	7.060	11.734	560.95	4.675
125.00	15.086	921.38	7.371	12.340	621.14	4.969
130.00	15.572	996.03	7.677	12.941	684.34	5.264
135.00	16.047	1077.1	7.978	13.538	750.54	5.560
140.00	16.496	1158.5	8.275	14.130	819.71	5.855
145.00	16.925	1242.0	8.566	14.716	891.83	6.151
150.00	17.358	1321.7	8.851	15.297	966.87	6.446
155.00	17.800	1415.6	9.133	15.874	1044.8	6.741
160.00	18.240	1505.7	9.411	16.446	1125.6	7.035
165.00	18.671	1590.0	9.685	17.014	1209.2	7.329
170.00	19.090	1692.4	9.955	17.577	1295.7	7.622
175.00	19.498	1788.9	10.222	18.136	1385.0	7.914
180.00	19.895	1887.4	10.485	18.691	1477.1	8.206
185.00	20.288	1987.8	10.745	19.242	1571.9	8.497
190.00	20.679	2099.2	11.001	19.788	1669.5	8.787
195.00	21.070	2194.6	11.254	20.330	1765.8	9.076
200.00	21.450	2300.9	11.505	20.869	1872.8	9.364
205.00	21.847	2405.2	11.752	21.403	1978.5	9.651
210.00	22.233	2514.4	11.997	21.934	2086.8	9.937
215.00	22.619	2631.5	12.240	22.462	2197.8	10.222
220.00	23.008	2745.6	12.480	22.986	2311.4	10.506
225.00	23.404	2861.5	12.718	23.508	2427.7	10.790
230.00	23.807	2974.6	12.955	24.027	2546.5	11.072
235.00	24.216	3099.7	13.190	24.543	2667.9	11.353
240.00	24.624	3221.8	13.424	25.057	2791.9	11.633
245.00	25.025	3345.9	13.657	25.569	2918.5	11.912
250.00	25.418	3472.0	13.888	26.079	3047.6	12.190
255.00	25.807	3600.1	14.118	26.586	3179.3	12.468
260.00	26.197	3730.1	14.347	27.091	3313.5	12.744
265.00	26.595	3862.1	14.574	27.593	3450.2	13.020
270.00	27.001	3996.1	14.800	28.094	3589.4	13.294
273.15	27.259	4081.5	14.942	28.409	3678.4	13.467
275.00	27.411	4132.1	15.026	28.594	3731.1	13.568
280.00	27.819	4270.2	15.251	29.091	3875.3	13.840
285.00	28.218	4410.3	15.475	29.587	4022.0	14.112
290.00	28.606	4552.3	15.698	30.081	4171.2	14.383
295.00	28.984	4696.3	15.920	30.573	4322.8	14.654
298.15	29.219	4788.0	16.059	30.883	4419.6	14.824
300.00	29.357	4842.2	16.141	31.064	4476.9	14.923

$H_{T-H_0^C}$ IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Hutchens, J. O., Cole, A. G. and Stout, J. W.,
Heat capacities from 11 to 305°K, and entropies of L-alanine and glycine,
J. Am. Chem. Soc. 82, 4813-4815 (1960).

TABLE 3
MOLAL THERMODYNAMIC FUNCTIONS FOR L-VALINE
 $((\text{CH}_3)_2\text{CH}(\text{NH}_2)\text{CHCOOH})$
SOLID PHASE

GRAM MOLECULAR WT. = 117.14892 GRAMS 1 CAL = 4.1840 JOULES
T DEG K = 273.15 + T DEG C

T DEG K	C_p^C CAL/DEG	$(H_T^0 - H_0^C)$ CAL	$(H_T^0 - H_0^C)/T$ CAL/DEG	S_T^0 CAL/DEG	$-(G_T^0 - H_0^C)$ CAL	$-(G_T^0 - H_0^C)/T$ CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.031	0.030	0.008	0.010	0.013	0.003
10.00	0.061	0.062	0.013	0.024	0.020	0.002
15.00	0.072	0.046	0.020	0.023	0.044	0.070
20.00	0.080	0.030	0.014	0.020	0.068	0.158
25.00	0.056	0.012	0.005	0.005	0.030	0.220
30.00	0.049	0.004	0.002	0.002	0.005	0.453
35.00	0.042	0.004	0.002	0.002	0.007	0.673
40.00	0.038	0.003	0.002	0.002	0.008	0.914
45.00	0.049	0.005	0.002	0.002	0.009	1.184
50.00	0.027	0.014	0.008	0.004	0.006	1.476
55.00	0.019	0.012	0.009	0.017	0.012	1.768
60.00	0.026	0.009	0.003	0.016	0.010	2.117
65.00	0.013	0.006	0.003	0.012	0.005	2.459
70.00	0.020	0.004	0.003	0.009	0.006	2.813
75.00	0.029	0.002	0.002	0.007	0.004	3.177
80.00	0.016	0.004	0.005	0.004	0.001	3.549
85.00	0.019	0.007	0.008	0.015	0.003	3.927
90.00	0.016	0.006	0.003	0.017	0.005	4.312
95.00	0.012	0.004	0.003	0.013	0.003	4.701
100.00	0.011	0.001	0.001	0.008	0.001	5.094
105.00	0.009	0.001	0.001	0.007	0.001	5.490
110.00	0.009	0.001	0.001	0.006	0.001	5.884
115.00	0.007	0.001	0.001	0.003	0.001	6.290
120.00	0.009	0.001	0.001	0.005	0.001	6.692
125.00	0.007	0.001	0.001	0.003	0.001	7.095
130.00	0.008	0.001	0.001	0.004	0.001	7.499
135.00	0.009	0.001	0.001	0.005	0.001	7.893
140.00	0.008	0.001	0.001	0.004	0.001	8.308
145.00	0.006	0.001	0.001	0.002	0.001	8.712
150.00	0.004	0.001	0.001	0.001	0.001	9.117
155.00	0.003	0.001	0.001	0.001	0.001	9.521
160.00	0.002	0.001	0.001	0.001	0.001	9.924
165.00	0.001	0.001	0.001	0.001	0.001	10.326
170.00	0.001	0.001	0.001	0.001	0.001	10.728
175.00	0.001	0.001	0.001	0.001	0.001	11.129
180.00	0.001	0.001	0.001	0.001	0.001	11.529
185.00	0.001	0.001	0.001	0.001	0.001	11.928
190.00	0.001	0.001	0.001	0.001	0.001	12.326
195.00	0.001	0.001	0.001	0.001	0.001	12.722
200.00	0.001	0.001	0.001	0.001	0.001	13.118
205.00	0.001	0.001	0.001	0.001	0.001	13.512
210.00	0.001	0.001	0.001	0.001	0.001	13.905
215.00	0.001	0.001	0.001	0.001	0.001	14.297
220.00	0.001	0.001	0.001	0.001	0.001	14.687
225.00	0.001	0.001	0.001	0.001	0.001	15.076
230.00	0.001	0.001	0.001	0.001	0.001	15.464
235.00	0.001	0.001	0.001	0.001	0.001	15.850
240.00	0.001	0.001	0.001	0.001	0.001	16.235
245.00	0.001	0.001	0.001	0.001	0.001	16.619
250.00	0.001	0.001	0.001	0.001	0.001	17.002
255.00	0.001	0.001	0.001	0.001	0.001	17.383
260.00	0.001	0.001	0.001	0.001	0.001	17.763
265.00	0.001	0.001	0.001	0.001	0.001	18.142
270.00	0.001	0.001	0.001	0.001	0.001	18.519
275.00	0.001	0.001	0.001	0.001	0.001	18.757
280.00	0.001	0.001	0.001	0.001	0.001	18.896
285.00	0.001	0.001	0.001	0.001	0.001	19.271
290.00	0.001	0.001	0.001	0.001	0.001	19.645
295.00	0.001	0.001	0.001	0.001	0.001	20.018
300.00	0.001	0.001	0.001	0.001	0.001	20.390
305.00	0.001	0.001	0.001	0.001	0.001	20.623
310.00	0.001	0.001	0.001	0.001	0.001	20.760

H_0^C IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Hutchens, J. O., Cole, A. G. and Stout, J. W.,
Heat capacities from 11 to 305 K, entropies, and free energies of formation
of l-valine, l-isoleucine, and l-leucine,
J. Phys. Chem. 67, 1128-1130 (1963).

TABLE 4

MOLAL THERMODYNAMIC FUNCTIONS FOR L-LEUCINE
 $(\text{CH}_3)_2\text{CHCH}_2(\text{NH}_2)\text{CHCOOH}$
SOLID PHASE

GRAM MOLECULAR WT. = 131.17601 GRAMS
T DEG K = 273.15 + T DEG C

1 CAL = 4.1840 JOULES

T DEG K	C _P CAL/DEG	(H _T ⁰ -H ₀ ^C) CAL	(H _T ⁰ -H ₀ ^C)/T CAL/DEG	S _T ⁰ CAL/DEG	-(C _T ⁰ -H ₀ ^C) CAL	-(G _T ⁰ -H ₀ ^C)/T CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.069	0.086	0.017	0.023	0.029	0.006
10.00	0.535	1.360	0.136	0.182	0.458	0.046
15.00	1.539	6.372	0.425	0.574	2.245	0.150
20.00	2.752	17.072	0.854	1.182	6.562	0.328
25.00	4.046	34.030	1.361	1.933	14.299	0.572
30.00	5.389	57.614	1.920	2.789	26.069	0.869
35.00	6.716	87.898	2.511	3.720	42.318	1.209
40.00	7.968	124.64	3.116	4.700	63.353	1.584
45.00	9.176	167.51	3.722	5.708	80.364	1.986
50.00	10.337	216.33	4.327	6.736	120.47	2.409
55.00	11.419	270.74	4.923	7.772	156.74	2.850
60.00	12.474	330.48	5.508	8.811	198.19	3.303
65.00	13.499	395.43	6.084	9.851	244.85	3.767
70.00	14.438	465.32	6.647	10.886	296.69	4.238
75.00	15.349	539.76	7.197	11.913	353.69	4.716
80.00	16.295	616.89	7.736	12.934	415.81	5.198
85.00	17.206	702.67	8.267	13.949	483.02	5.683
90.00	18.048	790.82	8.787	14.957	555.29	6.170
95.00	18.869	881.13	9.296	15.955	632.58	6.659
100.00	19.648	979.43	9.794	16.943	714.82	7.148
105.00	20.410	1079.6	10.282	17.920	801.98	7.638
110.00	21.167	1163.5	10.759	18.887	894.00	8.127
115.00	21.903	1291.2	11.228	19.844	990.83	8.616
120.00	22.618	1402.5	11.688	20.791	1092.4	9.104
125.00	23.323	1517.4	12.139	21.729	1198.7	9.590
130.00	24.015	1635.7	12.582	22.657	1309.7	10.075
135.00	24.689	1751.5	13.018	23.576	1425.3	10.558
140.00	25.356	1842.6	13.447	24.486	1545.4	11.039
145.00	26.020	2011.1	13.869	25.387	1670.1	11.518
150.00	26.674	2142.8	14.285	26.281	1799.3	11.995
155.00	27.318	2277.8	14.695	27.166	1932.9	12.470
160.00	27.960	2416.0	15.100	28.043	2070.9	12.943
165.00	28.608	2557.4	15.499	28.914	2213.3	13.414
170.00	29.261	2702.1	15.894	29.777	2360.1	13.883
175.00	29.910	2850.0	16.286	30.635	2511.1	14.349
180.00	30.544	3001.1	16.673	31.486	2666.4	14.813
185.00	31.160	3152.4	17.056	32.332	2826.0	15.275
190.00	31.767	3312.7	17.435	33.171	2985.7	15.735
195.00	32.378	3473.1	17.811	34.004	3157.7	16.193
200.00	33.003	3636.5	18.183	34.831	3329.7	16.649
205.00	33.644	3803.1	18.552	35.654	3506.0	17.102
210.00	34.296	3973.0	18.919	36.473	3686.3	17.554
215.00	34.950	4146.1	19.284	37.287	3870.7	18.003
220.00	35.608	4322.5	19.648	38.098	4059.1	18.451
225.00	36.270	4502.2	20.010	38.906	4251.7	18.996
230.00	36.900	4686.2	20.371	39.711	4449.2	19.240
235.00	37.670	4871.9	20.732	40.514	4648.8	19.782
240.00	38.401	5062.1	21.092	41.314	4852.3	20.222
245.00	39.127	5255.9	21.453	42.114	5061.9	20.661
250.00	39.858	5453.4	21.814	42.911	5274.5	21.098
255.00	40.598	5654.5	22.175	43.706	5491.0	21.533
260.00	41.336	5859.4	22.536	44.504	5711.5	21.967
265.00	42.157	6068.1	22.899	45.299	5936.0	22.400
270.00	42.942	6280.8	23.262	46.094	6164.5	22.832
275.00	43.467	6416.9	23.492	46.595	6310.5	23.103
280.00	43.780	6497.6	23.628	46.889	6397.0	23.262
285.00	44.656	6718.7	23.995	47.686	6631.4	23.691
290.00	45.577	6944.2	24.366	48.485	6873.9	24.119
295.00	46.539	7174.5	24.740	49.286	7118.3	24.546
298.15	47.597	7404.7	25.118	50.089	7366.7	24.972
298.15	48.146	7560.4	25.358	50.598	7525.3	25.240
300.00	48.504	7649.8	25.499	50.897	7619.2	25.397

H_T⁰^C IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Hutchens, J. O., Cole, A. G. and Stout, J. W.,
Heat capacities from 11 to 305 K., entropies, and free energies of formation
of L-valine, L-isoleucine, and L-leucine,
J. Phys. Chem. 67, 1128-1130 (1963).

TABLE 5

MOLAL THERMODYNAMIC FUNCTIONS FOR L-ISOLEUCINE
 $((\text{CH}_3)(\text{C}_2\text{H}_5)\text{CH}(\text{NH}_2)\text{CHCOOH})$
 SOLID PHASE

GRAM MOLECULAR WT. = 131.17601 GRAMS
 $T \text{ DEG K} = 273.15 + T \text{ DEG C}$

1 CAL = 4.1840 JOULES

T DEG K	C_p^{C} CAL/DEG	$(H_T^0 - H_0^{\text{C}})$ CAL	$(H_T^0 - H_0^{\text{C}})/T$ CAL/DEG	S_T^0 CAL/DEG	$-(Q_T^0 - Q_0^{\text{C}})$ CAL	$-(G_T^0 - G_0^{\text{C}})/T$ CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.185	0.100	0.022	0.029	0.037	0.007
10.00	0.397	1.542	0.154	0.215	0.557	0.056
15.00	1.519	6.745	0.452	0.623	2.566	0.171
20.00	2.636	16.943	0.850	1.203	7.075	0.354
25.00	3.756	32.864	1.315	1.907	14.805	0.592
30.00	4.876	54.807	1.827	2.704	26.293	0.877
35.00	6.002	82.448	2.370	3.569	41.054	1.199
40.00	7.126	117.25	2.931	4.483	62.066	1.552
45.00	8.257	157.57	3.502	5.431	86.841	1.930
50.00	9.382	202.79	4.076	6.404	116.42	2.328
55.00	10.517	255.72	4.650	7.393	150.91	2.744
60.00	12.657	315.20	5.220	8.393	190.37	3.173
65.00	13.793	376.06	5.786	9.399	234.65	3.613
70.00	14.928	444.12	6.345	10.407	284.36	4.062
75.00	16.064	517.04	6.894	11.413	338.91	4.519
80.00	16.200	594.82	7.435	12.416	398.48	4.981
85.00	17.327	677.53	7.971	13.419	463.07	5.448
90.00	17.463	764.91	8.499	14.417	532.67	5.919
95.00	18.598	856.52	9.016	15.408	607.23	6.392
100.00	19.733	952.18	9.522	16.389	686.73	6.867
105.00	20.868	1051.8	10.018	17.362	771.11	7.344
110.00	21.004	1155.9	10.505	18.326	860.27	7.821
115.00	21.896	1265.1	10.983	19.282	954.36	8.299
120.00	22.620	1374.4	11.453	20.230	1053.1	8.776
125.00	23.344	1483.6	11.914	21.167	1156.6	9.253
130.00	24.068	1607.6	12.366	22.096	1264.8	9.729
135.00	24.792	1722.6	12.812	23.016	1371.6	10.204
140.00	25.516	1835.0	13.250	13.928	1494.9	10.678
145.00	26.240	1948.7	13.681	14.831	1616.8	11.151
150.00	26.964	2115.7	14.105	15.726	1743.2	11.622
155.00	27.684	2251.7	14.523	16.614	1874.1	12.091
160.00	28.404	2381.6	14.935	17.493	2009.4	12.559
165.00	29.124	2511.5	15.341	18.365	2149.0	13.024
170.00	29.844	2616.2	15.742	19.230	2293.0	13.488
175.00	30.564	2624.1	16.138	20.088	2441.3	13.950
180.00	30.284	2717.2	16.529	20.939	2593.9	14.410
185.00	31.004	3127.3	16.915	31.784	2750.7	14.869
190.00	31.724	3248.4	17.297	32.622	2911.7	15.325
195.00	32.444	3446.4	17.674	33.453	3076.9	15.779
200.00	32.164	3607.4	18.047	34.278	3246.2	16.231
205.00	32.884	3775.2	18.416	35.097	3419.7	16.681
210.00	34.604	3943.0	18.781	35.910	3597.2	17.129
215.00	34.324	4115.7	19.143	36.719	3778.8	17.576
220.00	35.054	4295.5	19.502	37.522	3964.4	18.020
225.00	35.765	4468.3	19.859	38.321	4154.0	18.462
230.00	36.477	4649.2	20.214	39.116	4347.6	18.902
235.00	37.197	4833.1	20.566	39.907	4545.1	19.341
240.00	37.715	5020.1	20.917	40.695	4746.6	19.778
245.00	38.335	5210.2	21.266	41.479	4952.1	20.213
250.00	38.952	5405.5	21.614	42.259	5161.4	20.646
255.00	39.565	5599.7	21.960	43.037	5374.7	21.077
260.00	40.176	5799.1	22.304	43.811	5591.8	21.507
265.00	40.790	6001.5	22.647	44.582	5812.8	21.935
270.00	41.409	6207.0	22.989	45.350	6037.6	22.361
275.15	41.812	6338.1	23.204	45.833	6181.2	22.629
275.00	42.035	6415.6	23.330	46.116	6266.3	22.786
280.00	42.669	6627.4	23.669	46.879	6498.7	23.210
285.00	43.313	6842.3	24.008	47.640	6735.0	23.632
290.00	43.967	7060.5	24.347	48.399	6975.1	24.052
295.00	44.620	7282.0	24.685	49.156	7219.0	24.471
298.15	45.050	7423.3	24.898	49.632	7374.6	24.735
300.00	45.295	7506.8	25.023	49.912	7466.7	24.889

H_0^{C} IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Hutchens, J. O., Cole, A. G. and Stout, J. W.,
 Heat capacities from 11 to 305 K., entropies, and free energies of formation
 of l-valine, l-isoleucine, and l-leucine,
 J. Phys. Chem. 67, 1128-1130 (1963).

TABLE 6

MOLAL THERMODYNAMIC FUNCTIONS FOR L-TYROSINE
 $(\text{HOCH}_2\text{CH}_2(\text{NH}_2)\text{COOH})$
 SOLID PHASE

GRAM MOLECULAR WT. = 181.19292 GRAMS				1 CAL = 4,1840 JOULES		
T DEG K = 273.15 + T DEG C						
T	C_p°	$(H_{T=0}^0 - H_0^0)$	$(H_{T=0}^0 - H_0^0)/T$	S_T^0	$-(G_T^0 - G_0^0)$	$-(G_T^0 - G_0^0)/T$
DEG K	CAL/DEG	CAL	CAL/DEG	CAL/DEG	CAL	CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.035	0.044	0.009	0.012	0.015	0.003
10.00	0.070	0.097	0.070	0.093	0.232	0.023
15.00	0.093	0.303	0.234	0.312	1.175	0.078
20.00	0.054	10.336	0.517	0.699	3.634	0.182
25.00	0.000	22.386	0.895	1.232	8.402	0.336
30.00	4.323	40.655	1.355	1.894	16.166	0.539
35.00	5.712	67.734	1.878	2.664	27.523	0.786
40.00	7.079	97.732	2.443	3.517	42.948	1.074
45.00	8.413	136.47	3.033	4.428	62.789	1.395
50.00	9.705	181.80	3.636	5.382	87.299	1.746
55.00	10.880	233.30	4.242	6.363	116.65	2.121
60.00	12.117	290.55	4.842	7.358	150.95	2.516
65.00	13.107	353.39	5.437	8.364	190.25	2.927
70.00	14.120	421.49	6.021	9.373	234.59	3.351
75.00	15.112	494.57	6.594	10.381	283.98	3.786
80.00	16.093	572.59	7.157	11.387	338.40	4.230
85.00	17.065	652.50	7.712	12.392	397.85	4.681
90.00	17.959	743.09	8.257	13.393	462.51	5.137
95.00	18.766	834.92	8.789	14.386	531.77	5.598
100.00	19.561	930.74	9.307	15.369	606.16	6.062
105.00	20.391	1030.6	9.815	16.343	687.44	6.528
110.00	21.230	1134.7	10.315	17.311	769.58	6.996
115.00	22.058	1242.9	10.808	18.273	858.54	7.466
120.00	22.860	1355.2	11.293	19.229	952.30	7.936
125.00	23.630	1471.4	11.771	20.178	1050.8	8.407
130.00	24.419	1591.5	12.243	21.120	1154.1	8.877
135.00	25.239	1715.7	12.709	22.057	1264.0	9.348
140.00	26.041	1843.9	13.171	22.989	1374.6	9.819
145.00	26.815	1975.0	13.628	23.517	1491.9	10.289
150.00	27.593	2112.0	14.080	24.839	1613.8	10.759
155.00	28.391	2252.0	14.529	25.757	1740.3	11.228
160.00	29.204	2396.0	14.975	26.671	1871.4	11.696
165.00	30.021	2544.0	15.418	27.582	2007.0	12.164
170.00	30.834	2696.2	15.860	28.490	2147.2	12.630
175.00	31.638	2852.4	16.299	29.396	2291.9	13.096
180.00	32.431	3012.5	16.736	30.298	2441.1	13.562
185.00	33.222	3176.7	17.171	31.197	2594.9	14.026
190.00	34.017	3344.8	17.604	32.094	2753.1	14.490
195.00	34.818	3516.8	18.035	32.988	2915.8	14.953
200.00	35.626	3693.6	18.465	33.880	3087.0	15.415
205.00	36.440	3873.1	18.893	34.769	3254.6	15.876
210.00	37.261	4051.4	19.321	35.657	3430.7	16.336
215.00	38.085	4245.7	19.748	36.544	3611.2	16.796
220.00	38.904	4438.2	20.174	37.429	3796.1	17.255
225.00	39.713	4634.8	20.599	38.312	3985.4	17.713
230.00	40.512	4835.3	21.023	39.194	4179.2	18.170
235.00	41.307	5039.9	21.446	40.073	4377.4	18.627
240.00	42.107	5248.4	21.868	40.951	4579.9	19.083
245.00	42.918	5461.0	22.290	41.828	4786.9	19.538
250.00	43.741	5677.6	22.710	42.703	4998.2	19.993
255.00	44.575	5898.4	23.131	43.578	5213.9	20.447
260.00	45.416	6123.4	23.551	44.451	5434.0	20.900
265.00	46.258	6352.5	23.972	45.324	5658.4	21.353
270.00	47.098	6585.9	24.392	46.197	5887.2	21.805
275.00	47.624	6735.1	24.657	46.746	6033.6	22.089
280.00	47.932	6823.5	24.813	47.069	6120.4	22.256
285.00	48.762	7065.2	25.233	47.940	6357.9	22.707
290.00	49.591	7311.1	25.653	48.810	6599.3	23.157
295.00	50.423	7561.2	26.073	49.680	6846.0	23.607
298.15	51.257	7815.4	26.493	50.549	7096.5	24.056
298.15	51.782	7977.6	26.757	51.096	7256.7	24.339
300.00	52.188	8074.7	26.912	51.417	7351.5	24.505

H_0^0 IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Huffman, H. M. and Ellis, E. L.,
 Thermal data. VIII. The heat capacities, entropies and free energies of
 some amino acids,
 J. Am. Chem. Soc. 59, 2150-2152 (1937).

Cole, A. G., Hutchens, J. O. and Stout, J. W.,
 Heat capacities from 11 to 305 K. and entropies of l-phenylalanine,
 l-proline, l-tryptophane, and l-tyrosine. Some free energies of formation,
 J. Phys. Chem. 67, 1852-1855 (1963).

TABLE 7

MOLAL THERMODYNAMIC FUNCTIONS FOR L-PHENYLALANINE
 $(C_6H_5CH_2(NH_2)CHCOOH)$
 SOLID PHASE

GRAM MOLECULAR WT. = 165.19352 GRAMS
 T DEG K = 273.15 + T DEG C 1 CAL = 4.1840 JOULES

T DEG K	C _P CAL/DEG	(H _T ⁰ -H ₀ ^C) CAL	(H _T ⁰ -H ₀ ^C)/T CAL/DEG	S _T ⁰ CAL/DEG	-(Q _T ⁰ -H ₀ ^C) CAL	-(Q _T ⁰ -H ₀ ^C)/T CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.074	0.093	0.019	0.025	0.031	0.006
10.00	0.575	1.469	0.147	0.196	0.493	0.049
15.00	1.597	6.745	0.450	0.610	2.456	0.160
20.00	2.836	17.774	0.889	1.236	6.948	0.347
25.00	4.157	35.240	1.410	2.010	15.013	0.601
30.00	5.534	54.442	1.981	2.689	27.222	0.907
35.00	6.915	90.591	2.588	3.846	44.033	1.258
40.00	8.211	128.44	3.211	4.855	65.771	1.644
45.00	9.426	172.57	3.835	5.893	92.634	2.059
50.00	10.567	222.57	4.451	6.946	124.73	2.495
55.00	11.652	278.15	5.057	8.005	162.10	2.947
60.00	12.667	338.96	5.649	9.062	204.77	3.413
65.00	13.617	404.67	6.226	10.114	252.72	3.888
70.00	14.511	475.03	6.786	11.156	309.90	4.370
75.00	15.366	549.73	7.330	12.186	364.26	4.857
80.00	16.251	628.78	7.860	13.206	427.74	5.347
85.00	17.083	712.14	8.378	14.217	496.31	5.839
90.00	17.852	799.49	8.883	15.215	567.89	6.332
95.00	19.588	890.60	9.375	16.200	646.44	6.826
100.00	19.314	989.36	9.854	17.172	731.87	7.319
105.00	20.050	1083.8	10.322	18.132	820.14	7.811
110.00	20.778	1185.8	10.780	19.082	913.18	8.302
115.00	21.486	1291.5	11.231	20.021	1010.9	8.791
120.00	22.163	1400.7	11.672	20.950	1113.4	9.278
125.00	22.844	1513.2	12.105	21.869	1226.4	9.763
130.00	23.544	1629.1	12.532	22.778	1332.1	10.247
135.00	24.256	1748.6	12.953	23.680	1448.2	10.727
140.00	24.949	1871.7	13.369	24.575	1566.8	11.206
145.00	25.627	1998.1	13.780	25.462	1693.9	11.682
150.00	26.319	2128.0	14.186	26.343	1823.5	12.156
155.00	27.031	2261.3	14.589	27.217	1957.4	12.628
160.00	27.744	2398.3	14.989	28.087	2099.6	13.098
165.00	28.456	2538.8	15.387	28.952	2238.2	13.565
170.00	29.164	2682.8	15.781	29.812	2385.1	14.030
175.00	29.872	2830.4	16.174	30.667	2536.3	14.493
180.00	30.579	2981.6	16.564	31.519	2691.8	14.954
185.00	31.287	3136.2	16.953	32.366	2851.5	15.414
190.00	31.999	3294.4	17.339	33.210	3015.4	15.871
195.00	32.714	3456.2	17.724	34.060	3183.6	16.326
200.00	33.427	3621.6	18.108	34.888	3355.9	16.780
205.00	34.141	3790.5	18.490	35.722	3532.5	17.232
210.00	34.861	3963.0	18.871	36.553	3713.2	17.682
215.00	35.597	4139.1	19.252	37.382	3898.0	18.130
220.00	36.349	4319.0	19.632	38.209	4087.0	18.577
225.00	37.116	4502.6	20.012	39.034	4280.1	19.023
230.00	37.888	4690.1	20.392	39.859	4477.3	19.467
235.00	38.662	4881.5	20.772	40.682	4678.7	19.909
240.00	39.435	5076.8	21.153	41.504	4884.1	20.351
245.00	40.206	5275.9	21.534	42.325	5093.7	20.791
250.00	40.979	5478.8	21.915	43.145	5307.4	21.229
255.00	41.755	5685.7	22.297	43.964	5529.1	21.667
260.00	42.536	5896.4	22.678	44.782	5747.0	22.104
265.00	43.320	6111.0	23.060	45.600	5973.0	22.539
270.00	44.106	6329.6	23.443	46.417	6203.0	22.974
275.00	44.860	6469.3	23.824	46.931	6350.0	23.427
280.00	44.890	6552.1	23.826	47.234	6437.1	23.408
285.00	45.671	6778.5	24.209	48.049	6675.3	23.841
290.00	46.448	7008.8	24.592	48.865	6917.6	24.272
295.00	47.221	7242.9	24.976	49.679	7164.0	24.703
298.15	48.481	7481.0	25.359	50.493	7414.4	25.134
300.00	48.768	7722.9	25.743	51.306	7574.3	25.404
					7668.9	25.563

H₀^C IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Cole, A. G., Hutchens, J. O. and Stout, J. W.,
 Heat capacities from 11 to 305 K. and entropies of L-phenylalanine,
 L-proline, L-tryptophane, and L-tyrosine. Some free energies of formation,
J. Phys. Chem. 67, 1852-1855 (1963).

TABLE 8

MOLAL THERMODYNAMIC FUNCTIONS FOR L-TRYPTOPHANE
 $(C_8H_6NCH_2(NH_2)CHCOOH)$
 SOLID PHASE

GRAM MOLECULAR WT. = 204.23049 GRAMS
 T DEG K = 273.15 + T DEG C

1 CAL = 4.1840 JOULES

T	C_p^c	$(H_T^0 - H_0^c)$	$(H_T^0 - H_0^c)/T$	S_T^0	$-(G_T^0 - H_0^c)$	$-(G_T^0 - H_0^c)/T$
DEG K	CAL/DEG	CAL	CAL/DEG	CAL/DEG	CAL	CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.116	0.144	0.029	0.038	0.048	0.010
10.00	0.896	2.293	0.229	0.306	0.770	0.077
15.00	2.393	10.371	0.691	0.941	3.740	0.249
20.00	3.947	26.248	1.312	1.843	10.622	0.531
25.00	5.495	49.850	1.994	2.890	22.410	0.896
30.00	6.995	81.102	2.703	4.026	34.673	1.322
35.00	8.434	119.72	3.420	5.213	62.755	1.793
40.00	9.742	165.21	4.130	6.426	91.849	2.296
45.00	10.991	217.04	4.823	7.646	127.03	2.823
50.00	12.200	275.06	5.501	8.867	168.31	3.366
55.00	13.310	338.86	6.161	10.083	215.69	3.922
60.00	14.412	408.14	6.802	11.288	269.12	4.485
65.00	15.459	482.82	7.428	12.483	328.55	5.055
70.00	16.429	562.57	8.037	13.664	393.93	5.628
75.00	17.390	647.12	8.628	14.831	465.17	6.202
80.00	18.383	736.55	9.207	15.985	542.21	6.778
85.00	19.328	830.85	9.775	17.128	625.00	7.353
90.00	20.212	929.72	10.330	18.258	713.47	7.927
95.00	21.092	1033.0	10.873	19.374	807.55	8.501
100.00	21.973	1140.7	11.407	20.478	907.19	9.072
105.00	22.838	1252.7	11.931	21.572	1012.3	9.641
110.00	23.674	1369.0	12.445	22.652	1122.9	10.208
115.00	24.514	1489.5	12.952	23.724	1238.8	10.772
120.00	25.341	1614.1	13.451	24.785	1360.1	11.334
125.00	26.169	1742.9	13.943	25.836	1486.7	11.893
130.00	27.025	1875.8	14.430	26.879	1618.5	12.450
135.00	27.898	2013.2	14.912	27.916	1755.5	13.003
140.00	28.763	2154.8	15.392	28.946	1897.6	13.554
145.00	29.606	2300.7	15.867	29.970	2044.9	14.103
150.00	30.444	2450.9	16.339	30.988	2197.3	14.649
155.00	31.296	2605.2	16.808	32.000	2354.8	15.192
160.00	32.159	2763.8	17.274	33.007	2517.3	15.733
165.00	33.019	2926.8	17.738	34.010	2684.8	16.272
170.00	33.871	3094.0	18.200	35.008	2857.4	16.808
175.00	34.721	3265.5	18.660	36.002	3034.9	17.342
180.00	35.576	3441.2	19.118	36.992	3217.4	17.874
185.00	36.442	3621.3	19.575	37.979	3404.8	18.404
190.00	37.316	3805.7	20.030	38.962	3597.2	18.933
195.00	38.191	3994.4	20.484	39.943	3794.4	19.459
200.00	39.064	4187.6	20.938	40.921	3996.6	19.983
205.00	39.942	4385.1	21.391	41.896	4203.6	20.506
210.00	40.825	4587.0	21.843	42.869	4415.6	21.026
215.00	41.713	4792.4	22.295	43.840	4632.3	21.546
220.00	42.601	5004.1	22.746	44.810	4854.0	22.063
225.00	43.489	5219.4	23.197	45.777	5080.4	22.580
230.00	44.382	5439.0	23.548	46.742	5311.7	23.094
235.00	45.287	5663.2	24.099	47.707	5547.9	23.608
240.00	46.200	5891.9	24.550	48.670	5788.8	24.120
245.00	47.120	6125.2	25.001	49.632	6034.5	24.631
250.00	48.042	6363.1	25.453	50.593	6285.1	25.140
255.00	48.964	6605.6	25.904	51.553	6540.5	25.649
260.00	49.884	6852.8	26.357	52.513	6800.6	26.156
265.00	50.800	7104.5	26.809	53.472	7065.6	26.663
270.00	51.712	7360.8	27.262	54.430	7335.4	27.168
275.15	52.625	7524.6	27.547	55.033	7507.8	27.486
275.00	52.623	7621.6	27.715	55.387	7609.9	27.672
280.00	53.540	7887.0	28.168	56.344	7889.2	28.176
285.00	54.468	8157.0	28.621	57.299	8173.3	28.678
290.00	55.409	8431.7	29.075	58.255	8462.2	29.180
295.00	56.360	8711.1	29.529	59.210	8755.9	29.681
298.15	56.958	8889.6	29.816	59.812	8943.3	29.996
300.00	57.307	8995.4	29.954	60.165	9054.3	30.181

H_0^c IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Cole, A. G., Hutchens, J. O. and Stout, J. W.,
 Heat capacities from 11 to 305°K. and entropies of l-phenylalanine,
 l-proline, l-tryptophane, and l-tyrosine. Some free energies of formation,
 J. Phys. Chem. 67, 1852-1855 (1963).

TABLE 9

MOLAL THERMODYNAMIC FUNCTIONS FOR L-ASPARTIC ACID
 $(\text{HOOCCH}_2(\text{NH}_2)\text{CHCOOH})$

SOLID PHASE

GRAM MOLECULAR WT. = 133.10469 GRAMS
 $T \text{ DEG K} = 273.15 + T \text{ DEG C}$

1 CAL = 4.1840 JOULES

T DEG K	C_p^C CAL/DEG	$(H_T^0 - H_0^C)$ CAL	$(H_T^0 - H_0^C)/T$ CAL/DEG	S_T^0 CAL/DEG	$-(Q_T^0 - H_0^C)$ CAL	$-(Q_T^0 - H_0^C)/T$ CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.022	0.022	0.004	0.007	0.000	0.002
10.00	0.100	0.100	0.024	0.059	0.145	0.015
15.00	0.199	2.324	0.156	0.206	0.757	0.050
20.00	0.444	7.424	0.371	0.493	2.444	0.122
25.00	0.467	17.131	0.665	0.922	5.928	0.237
30.00	3.619	32.310	1.077	1.473	11.870	0.396
35.00	4.823	53.405	1.526	2.121	20.818	0.595
40.00	6.017	80.516	2.013	2.843	33.200	0.830
45.00	7.184	113.53	2.523	3.619	49.336	1.096
50.00	8.294	152.26	3.045	4.434	69.457	1.389
55.00	9.340	196.36	3.570	5.274	93.720	1.704
60.00	10.343	245.59	4.073	6.130	122.23	2.037
65.00	11.288	299.69	4.611	6.996	155.04	2.385
70.00	12.154	358.34	5.117	7.865	192.19	2.746
75.00	12.962	421.19	5.616	8.732	233.68	3.116
80.00	13.796	488.14	6.102	9.596	279.20	3.494
85.00	14.578	559.10	6.578	10.456	329.63	3.878
90.00	15.297	633.61	7.042	11.310	384.05	4.267
95.00	15.959	711.97	7.494	12.155	442.71	4.660
100.00	16.601	793.38	7.944	12.990	505.58	5.056
105.00	17.242	877.94	8.361	13.815	572.59	5.453
110.00	17.852	965.58	8.778	14.630	643.71	5.852
115.00	18.432	1056.2	9.185	15.436	718.87	6.251
120.00	19.011	1149.9	9.582	16.233	798.05	6.650
125.00	19.577	1246.5	9.970	17.020	881.18	7.049
130.00	20.124	1345.6	10.350	17.798	968.23	7.448
135.00	20.682	1447.6	10.723	18.568	1059.2	7.846
140.00	21.219	1552.3	11.088	19.330	1153.9	8.242
145.00	21.747	1657.0	11.447	20.084	1252.4	8.638
150.00	22.273	1765.8	11.799	20.830	1354.7	9.032
155.00	22.795	1882.5	12.145	21.569	1460.7	9.424
160.00	23.321	1997.8	12.496	22.301	1570.4	9.815
165.00	23.842	2115.7	12.822	23.027	1683.7	10.204
170.00	24.350	2236.2	13.154	23.746	1800.7	10.592
175.00	24.842	2355.2	13.481	24.459	1921.2	10.978
180.00	25.334	2484.6	13.804	25.166	2045.9	11.363
185.00	25.854	2612.6	14.122	25.867	2172.8	11.745
190.00	26.378	2743.2	14.438	26.564	2303.9	12.126
195.00	26.864	2870.4	14.751	27.256	2438.9	12.505
200.00	27.351	3012.0	15.060	27.942	2576.9	12.882
205.00	27.823	3149.7	15.366	28.624	2717.9	13.258
210.00	28.291	3290.2	15.668	29.300	2862.7	13.632
215.00	28.772	3432.9	15.967	29.971	3010.9	14.004
220.00	29.265	3570.0	16.263	30.638	3162.4	14.375
225.00	29.752	3725.5	16.558	31.301	3317.2	14.743
230.00	30.262	3875.6	16.850	31.961	3475.4	15.110
235.00	30.763	4026.2	17.141	32.617	3636.8	15.476
240.00	31.264	4183.2	17.430	33.270	3801.6	15.840
245.00	31.761	4340.8	17.718	33.920	3969.5	16.202
250.00	32.259	4500.9	18.003	34.566	4140.8	16.563
255.00	32.755	4663.4	18.288	35.210	4315.2	16.922
260.00	33.248	4826.4	18.571	35.851	4492.9	17.280
265.00	33.741	4995.9	18.852	36.489	4673.7	17.637
270.00	34.233	5165.8	19.133	37.124	4857.7	17.992
275.00	34.743	5274.1	19.309	37.523	4975.3	18.215
280.00	34.725	5338.2	19.412	37.757	5044.9	18.345
285.00	35.223	5513.1	19.690	38.387	5235.3	18.698
290.00	35.730	5690.4	19.966	39.015	5428.8	19.048
295.00	36.248	5870.4	20.243	39.641	5625.4	19.398
300.00	36.775	6052.9	20.518	40.265	5825.2	19.746
305.00	37.108	6169.3	20.692	40.657	5952.7	19.965
310.00	37.304	6238.1	20.794	40.887	6028.1	20.094

H_0^C IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Huffman, H. M. and Borsook, H.,
 Thermal data. I. The heat capacities, entropies and free energies of
 seven organic compounds containing nitrogen,
J. Am. Chem. Soc. 54, 4297-4301 (1932).

Hutchens, J. O., Cole, A. G., Robbie, R. A. and Stout, J. W.,
 Heat capacities from 11 to 305°K, entropies and free energies of formation
 of l-asparagine monohydrate, l-aspartic acid, l-glutamic acid, and l-glutamine,
J. Biol. Chem. 238, 2407-2412 (1963).

TABLE 10

MOLAL THERMODYNAMIC FUNCTIONS FOR L-ASPARAGINE MONOHYDRATE
 $(\text{NH}_2\text{COCH}_2(\text{NH}_2)\text{CHCOOH}\cdot\text{H}_2\text{O})$
 SOLID PHASE

GRAM MOLECULAR WT. = 150.13530 GRAMS
 $T \text{ DEG K} = 273.15 + T \text{ DEG C}$

T DEG K	C_p° CAL/DEG	$(H_{T=0}^0 - H_0^0)$ CAL	$(H_{T=0}^0 - H_0^0)/T$ CAL/DEG	S_T^0 CAL/DEG	$-(G_{T=0}^0 - G_0^0)$ CAL	$-(G_{T=0}^0 - G_0^0)/T$ CAL/DEG	
0.00	0.000	0.000	0.000	0.000	0.000	0.000	
5.00	0.020	0.020	0.005	0.007	0.008	0.002	
10.00	0.164	0.409	0.041	0.054	0.135	0.014	
15.00	0.262	0.693	0.140	0.186	0.692	0.046	
20.00	0.354	0.980	0.329	0.439	2.197	0.110	
25.00	0.429	1.262	0.615	0.626	3.307	0.212	
30.00	0.497	1.545	0.991	1.348	10.642	0.356	
35.00	0.550	1.830	1.445	1.987	16.985	0.542	
40.00	0.598	2.131	1.958	2.727	30.732	0.768	
45.00	0.638	2.417	2.515	3.546	46.383	1.031	
50.00	0.672	2.697	3.101	4.427	66.222	1.326	
55.00	0.700	2.977	3.705	5.355	90.732	1.650	
60.00	0.719	3.253	4.317	6.315	119.89	1.998	
65.00	0.746	3.538	4.933	7.302	153.93	2.368	
70.00	0.771	3.823	5.547	8.304	192.94	2.776	
75.00	0.794	4.103	6.153	9.313	225.48	3.160	
80.00	0.816	4.383	6.752	10.328	258.03	3.576	
85.00	0.832	4.660	7.342	11.345	340.26	4.003	
90.00	0.847	4.937	7.924	12.363	339.53	4.439	
95.00	0.861	5.212	8.493	13.376	461.88	4.883	
100.00	0.875	5.492	9.049	14.382	533.28	5.333	
105.00	0.887	5.767	9.594	15.382	607.69	5.788	
110.00	0.901	6.042	10.129	16.375	667.09	6.246	
115.00	0.914	6.318	10.654	17.362	711.43	6.708	
120.00	0.927	6.593	11.249	18.343	860.70	7.172	
125.00	0.940	6.868	11.837	11.678	19.316	7.639	
130.00	0.952	7.143	12.427	20.283	1055.8	8.107	
135.00	0.964	7.419	12.969	21.244	1157.7	8.575	
140.00	0.976	7.694	13.154	22.199	1265.3	9.045	
145.00	0.987	7.969	13.633	23.147	1379.6	9.515	
150.00	0.997	8.244	2115.7	14.105	24.030	1497.7	9.985
155.00	0.997	8.519	2258.6	14.572	25.027	1620.5	10.455
160.00	0.997	8.794	2405.3	15.033	25.956	1748.0	10.925
165.00	0.997	9.069	2555.7	15.489	26.884	1860.1	11.395
170.00	0.997	9.344	2710.0	15.941	27.805	2016.8	11.864
175.00	0.997	9.619	2865.0	16.388	28.721	2158.1	12.332
180.00	0.997	9.894	3020.8	16.832	29.632	2304.0	12.800
185.00	0.997	10.168	3176.3	17.272	30.539	2454.5	13.267
190.00	0.997	10.443	3324.6	17.708	31.442	2609.4	13.734
195.00	0.997	10.718	3474.4	18.140	32.340	2768.9	14.199
200.00	0.997	11.003	3715.0	18.569	33.233	2932.8	14.664
205.00	0.997	11.277	3869.0	18.994	34.122	3101.2	15.128
210.00	0.997	11.552	4023.0	19.416	35.007	3274.0	15.591
215.00	0.997	11.826	4264.6	19.835	35.888	3451.3	16.052
220.00	0.997	12.101	4455.4	20.252	36.765	3632.9	16.513
225.00	0.997	12.376	4547.9	20.666	37.639	3818.9	16.973
230.00	0.997	12.651	4847.4	21.078	38.510	4009.3	17.432
235.00	0.997	12.926	5048.6	21.488	39.377	4204.0	17.889
240.00	0.997	13.199	5250.0	21.896	40.242	4403.0	18.346
245.00	0.997	13.474	5463.9	22.302	41.103	4606.4	18.802
250.00	0.997	13.844	5676.5	22.706	41.962	4814.1	19.256
255.00	0.997	14.114	5892.7	23.109	42.819	5026.0	19.710
260.00	0.997	14.382	6112.6	23.510	43.672	5242.3	20.163
265.00	0.997	14.652	6330.0	23.909	44.523	5462.7	20.614
270.00	0.997	14.922	6552.4	24.307	45.372	5687.5	21.065
275.00	0.997	15.192	6707.6	24.557	45.905	5831.2	21.348
280.00	0.997	15.461	6794.3	24.703	46.217	5916.5	21.514
285.00	0.997	15.730	7027.3	25.097	47.061	6149.6	21.963
290.00	0.997	16.000	7264.8	25.491	47.901	6387.1	22.411
295.00	0.997	16.269	7506.0	25.883	48.740	6628.7	22.857
298.15	0.997	16.313	7750.7	26.274	49.577	6874.5	23.303
300.00	0.997	16.763	7906.8	26.519	50.103	7031.4	23.584
300.00	0.997	16.924	7999.1	26.664	50.412	7124.4	23.748

H_0^0 IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Huffman, H. M. and Borscock, H.,
 Thermal data. I. The heat capacities, entropies and free energies of
 seven organic compounds containing nitrogen,
 J. Am. Chem. Soc. 54, 4297-4301 (1932).

Hutchens, J. O., Cole, A. G., Robie, R. A. and Stout, J. W.,
 Heat capacities from 11 to 305°K, entropies and free energies of formation
 of l-asparagine monohydrate, l-aspartic acid, l-glutamic acid, and l-glutamine,
 J. Biol. Chem. 238, 2407-2412 (1963).

TABLE 11

MOLAL THERMODYNAMIC FUNCTIONS FOR L-GLUTAMIC ACID
 $(HOOC(CH_2)_2(NH_2)CHCOOH)$
 SOLID PHASE

GRAM MOLECULAR WT. = 147.13178 GRAMS
 $T \text{ DEG K} = 273.15 + T \text{ DEG C}$

T DEG K	C_p^C CAL/DEG	$(H_T^0 - H_0^C)$ CAL	$(H_T^0 - H_0^C)/T$ CAL/DEG	S_T^0 CAL/DEG	$-(G_T^0 - H_0^C)$ CAL	$-(G_T^0 - H_0^C)/T$ CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.024	0.024	0.006	0.008	0.011	0.002
10.00	0.048	0.048	0.009	0.006	0.017	0.0017
15.00	0.065	0.065	0.015	0.018	0.043	0.006
20.00	0.094	0.094	0.021	0.024	0.060	0.0132
25.00	0.126	0.126	0.029	0.026	0.091	0.0252
30.00	0.158	0.158	0.037	0.032	0.125	0.0418
35.00	0.184	0.184	0.045	0.036	0.160	0.0629
40.00	0.211	0.211	0.053	0.042	0.196	0.0879
45.00	0.239	0.239	0.061	0.048	0.234	0.1163
50.00	0.262	0.262	0.069	0.054	0.274	0.1477
55.00	0.287	0.287	0.077	0.063	0.313	0.1816
60.00	0.310	0.310	0.085	0.072	0.352	0.2175
65.00	0.335	0.335	0.093	0.081	0.390	0.2552
70.00	0.358	0.358	0.101	0.090	0.429	0.2942
75.00	0.380	0.380	0.109	0.099	0.467	0.3344
80.00	0.407	0.407	0.117	0.108	0.506	0.3754
85.00	0.430	0.430	0.125	0.117	0.545	0.4172
90.00	0.451	0.451	0.133	0.126	0.584	0.4596
95.00	0.470	0.470	0.141	0.135	0.623	0.5024
100.00	0.487	0.487	0.149	0.144	0.662	0.5456
105.00	0.503	0.503	0.157	0.153	0.701	0.5890
110.00	0.517	0.517	0.165	0.162	0.737	0.6325
115.00	0.530	0.530	0.173	0.171	0.775	0.6762
120.00	0.541	0.541	0.181	0.180	0.813	0.7199
125.00	0.553	0.553	0.189	0.189	0.851	0.7636
130.00	0.563	0.563	0.197	0.198	0.889	0.8073
135.00	0.572	0.572	0.205	0.207	0.927	0.8509
140.00	0.580	0.580	0.213	0.212	0.962	0.8944
145.00	0.587	0.587	0.221	0.219	1.001	0.9379
150.00	0.593	0.593	0.228	0.226	1.038	0.9812
155.00	0.598	0.598	0.235	0.233	1.077	1.0244
160.00	0.602	0.602	0.242	0.241	1.116	1.0674
165.00	0.605	0.605	0.249	0.248	1.154	1.1103
170.00	0.607	0.607	0.255	0.254	1.190	1.1530
175.00	0.609	0.609	0.261	0.260	2.092	1.1955
180.00	0.611	0.611	0.267	0.266	2.228	1.2379
185.00	0.612	0.612	0.273	0.272	2.368	1.2801
190.00	0.613	0.613	0.278	0.277	2.512	1.3222
195.00	0.614	0.614	0.283	0.282	2.659	1.3640
200.00	0.615	0.615	0.288	0.287	2.811	1.4057
205.00	0.616	0.616	0.293	0.292	2.966	1.4473
210.00	0.616	0.616	0.298	0.297	3.126	1.4886
215.00	0.616	0.616	0.303	0.302	3.289	1.5298
220.00	0.616	0.616	0.308	0.307	3.455	1.5708
225.00	0.616	0.616	0.313	0.312	3.626	1.6117
230.00	0.616	0.616	0.318	0.317	3.800	1.6523
235.00	0.616	0.616	0.323	0.322	3.978	1.6929
240.00	0.616	0.616	0.328	0.327	4.159	1.7332
245.00	0.616	0.616	0.333	0.332	4.344	1.7734
250.00	0.616	0.616	0.338	0.337	4.533	1.8135
255.00	0.616	0.616	0.343	0.342	4.726	1.8534
260.00	0.616	0.616	0.348	0.347	4.922	1.8932
265.00	0.616	0.616	0.353	0.352	5.121	1.9328
270.00	0.616	0.616	0.358	0.357	5.325	1.9723
275.00	0.616	0.616	0.363	0.362	5.545	2.0171
280.00	0.616	0.616	0.368	0.367	5.761	2.0516
285.00	0.616	0.616	0.373	0.372	5.972	2.0908
290.00	0.616	0.616	0.378	0.377	6.183	2.1288
295.00	0.616	0.616	0.383	0.382	6.394	2.1676
300.00	0.616	0.616	0.388	0.387	6.605	2.2092

H_0^C IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Hutchens, J. O., Cole, A. G., Robie, R. A. and Stout, J. W.,
 Heat capacities from 11 to 305°K, entropies and free energies of formation
 of l-asparagine monohydrate, l-aspartic acid, l-glutamic acid, and l-glutamine,
J. Biol. Chem. 238, 2407-2412 (1963).

TABLE 12

MOLAL THERMODYNAMIC FUNCTIONS FOR L-GLUTAMINE
 $(\text{NH}_2\text{CO}(\text{CH}_2)_2(\text{NH}_2)\text{CHCOOH})$
 SOLID PHASE

GRAM MOLECULAR WT. = 146.14705 GRAMS

T DEG K = 273.15 + T DEG C

1 CAL = 4.1840 JOULES

T DEG K	c_p^C CAL/DEG	$(H_T^0 - H_0^C)$ CAL	$(H_T^0 - H_0^C)/T$ CAL/DEG	S_T^0 CAL/DEG	$-(\alpha_T^0 - \alpha_0^C)$ CAL	$-(\alpha_T^0 - \alpha_0^C)/T$ CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.022	0.028	0.006	0.007	0.009	0.002
10.00	0.177	0.444	0.044	0.059	0.148	0.015
15.00	0.598	2.244	0.150	0.200	0.749	0.050
20.00	1.389	7.048	0.352	0.471	2.363	0.118
25.00	2.507	16.686	0.667	0.896	5.715	0.229
30.00	3.811	32.420	1.081	1.466	11.564	0.385
35.00	5.217	54.969	1.571	2.159	20.580	0.588
40.00	6.617	84.568	2.114	2.947	33.310	0.833
45.00	7.991	121.10	2.691	3.806	50.166	1.115
50.00	9.304	164.38	3.288	4.717	71.455	1.429
55.00	10.527	213.99	3.891	5.661	97.389	1.771
60.00	11.693	269.56	4.493	6.628	128.10	2.135
65.00	12.771	330.77	5.089	7.607	163.69	2.518
70.00	13.760	397.13	5.673	8.590	204.18	2.917
75.00	14.695	468.28	6.244	9.572	249.59	3.328
80.00	15.615	544.05	6.801	10.549	299.89	3.749
85.00	16.554	624.49	7.347	11.524	355.08	4.177
90.00	17.391	709.40	7.882	12.495	415.13	4.613
95.00	18.125	798.21	8.402	13.455	480.00	5.053
100.00	18.863	890.68	8.907	14.403	549.65	5.497
105.00	19.593	986.82	9.398	15.341	624.02	5.943
110.00	20.316	1086.6	9.878	16.270	703.05	6.391
115.00	21.031	1190.0	10.348	17.189	786.70	6.841
120.00	21.691	1296.8	10.807	18.098	874.92	7.291
125.00	22.323	1406.8	11.255	18.996	967.66	7.741
130.00	22.991	1520.1	11.693	19.884	1064.9	8.191
135.00	23.678	1636.8	12.124	20.765	1166.5	8.641
140.00	24.339	1756.8	12.549	21.638	1272.5	9.089
145.00	24.977	1880.1	12.966	22.503	1382.9	9.537
150.00	25.615	2006.6	13.377	23.361	1497.5	9.983
155.00	26.259	2136.3	13.783	24.211	1616.5	10.429
160.00	26.903	2269.2	14.182	25.055	1739.6	10.873
165.00	27.546	2405.3	14.578	25.893	1867.0	11.315
170.00	28.183	2544.6	14.969	26.725	1998.5	11.756
175.00	28.815	2687.1	15.355	27.551	2134.2	12.196
180.00	29.440	2832.8	15.738	28.371	2274.0	12.634
185.00	30.061	2981.5	16.116	29.186	2417.9	13.070
190.00	30.679	3133.4	16.491	29.996	2565.9	13.505
195.00	31.296	3288.3	16.863	30.801	2717.9	13.938
200.00	31.916	3446.4	17.232	31.601	2873.9	14.369
205.00	32.539	3607.5	17.597	32.397	3033.9	14.799
210.00	33.163	3771.7	17.961	33.189	3197.9	15.228
215.00	33.787	3939.1	18.321	33.976	3365.8	15.655
220.00	34.413	4109.6	18.680	34.760	3537.6	16.080
225.00	35.049	4283.3	19.037	35.541	3713.4	16.504
230.00	35.696	4460.1	19.392	36.318	3893.0	16.926
235.00	36.351	4640.2	19.746	37.093	4076.5	17.347
240.00	37.007	4823.6	20.098	37.865	4263.9	17.766
245.00	37.654	5010.3	20.450	38.635	4455.2	18.184
250.00	38.285	5200.1	20.801	39.402	4650.3	18.601
255.00	38.897	5393.1	21.149	40.166	4849.2	19.016
260.00	39.492	5589.1	21.497	40.927	5051.9	19.431
265.00	40.075	5788.0	21.842	41.685	5258.5	19.843
270.00	40.655	5989.8	22.185	42.439	5468.8	20.255
273.15	41.021	6118.5	22.400	42.913	5603.2	20.513
275.00	41.237	6194.6	22.526	43.191	5682.9	20.665
280.00	41.823	6402.2	22.865	43.939	5900.7	21.074
285.00	42.408	6612.8	23.203	44.684	6122.2	21.482
290.00	42.987	6826.3	23.539	45.427	6347.5	21.888
295.00	43.563	7042.7	23.873	46.167	6576.5	22.293
298.15	43.927	7180.4	24.053	46.631	6722.7	22.548
300.00	44.144	7231.4	24.276	47.104	6862.2	22.697

 H_0^C IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Butchers, J. O., Cole, A. G., Robie, R. A. and Stout, J. W.,
 Heat capacities from 11 to 305°K, entropies and free energies of formation
 of l-asparagine monohydrate, l-aspartic acid, l-glutamic acid, and l-glutamine,
 J. Biol. Chem. 238, 2407-2412 (1963).

TABLE 13

MOLAL THERMODYNAMIC FUNCTIONS FOR L-LYSINE HYDROCHLORIDE
 $((\text{NH}_3^+ \text{Cl})(\text{CH}_2)_4(\text{NH}_2)\text{CHCOOH})$
 SOLID PHASE

GRAM MOLECULAR WT. = 182.65165 GRAMS			1 CAL = 4.1840 JOULES			
T DEG K = 273.15 + T DEG C						
T	C_p°	$(H_{T=0}^0 - H_0^0)$	$(H_{T=0}^0 - H_0^0)/T$	S_T^0	$-(G_{T=0}^0 - G_0^0)$	$-(G_{T=0}^0 - G_0^0)/T$
DEG K	CAL/DEG	CAL	CAL/DEG	CAL/DEG	CAL	CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.062	0.077	0.015	0.021	0.026	0.005
10.00	0.129	1.248	0.125	0.166	0.415	0.041
15.00	0.152	6.139	0.409	0.549	2.089	0.139
20.00	0.194	17.290	0.864	1.181	6.322	0.316
25.00	0.496	35.782	1.431	1.999	14.202	0.568
30.00	0.191	62.477	2.082	2.968	26.565	0.886
35.00	0.913	97.747	2.793	4.052	44.074	1.259
40.00	0.584	141.51	3.538	5.218	67.221	1.681
45.00	11.203	193.50	4.300	6.441	96.350	2.141
50.00	12.838	253.62	5.072	7.706	131.70	2.634
55.00	14.244	321.76	5.850	9.004	175.47	3.154
60.00	15.842	397.38	6.623	10.319	221.77	3.696
65.00	17.257	480.18	7.387	11.644	276.68	4.257
70.00	18.549	567.75	8.139	12.971	338.22	4.832
75.00	19.785	665.59	8.875	14.293	406.38	5.418
80.00	21.043	767.66	9.596	15.610	481.14	6.014
85.00	22.277	875.99	10.306	16.923	562.47	6.617
90.00	23.481	999.19	11.002	18.228	650.35	7.226
95.00	24.394	1109.6	11.680	19.520	744.73	7.939
100.00	25.471	1234.1	12.341	20.797	847.53	8.455
105.00	26.769	1363.6	12.987	22.060	952.67	9.073
110.00	27.373	1498.1	13.619	23.311	1066.1	9.692
115.00	28.314	1637.3	14.238	24.548	1185.8	10.311
120.00	29.213	1781.1	14.843	25.773	1311.6	10.930
125.00	30.077	1927.4	15.435	26.983	1443.5	11.548
130.00	30.828	2081.9	16.015	28.179	1581.4	12.164
135.00	31.773	2238.7	16.583	29.362	1725.2	12.779
140.00	32.551	2399.6	17.140	30.532	1875.0	13.393
145.00	33.350	2664.2	17.686	31.690	2030.5	14.004
150.00	34.150	2731.5	18.223	32.835	2141.4	14.612
155.00	34.950	2806.3	18.750	33.965	2258.0	15.210
160.00	35.730	3063.0	19.269	35.091	2331.5	15.822
165.00	36.452	3264.6	19.779	36.202	2709.8	16.423
170.00	37.250	3448.0	20.282	37.303	2893.5	17.021
175.00	38.002	3636.1	20.778	38.394	3082.8	17.616
180.00	38.742	3824.0	21.266	39.475	3277.4	18.208
185.00	39.462	4023.5	21.749	40.546	3477.5	18.797
190.00	40.191	4222.6	22.224	41.608	3682.9	19.384
195.00	40.911	4420.4	22.694	42.661	3893.6	19.967
200.00	41.635	4631.8	23.159	43.706	4104.5	20.547
205.00	42.367	4841.8	23.616	44.743	4320.6	21.125
210.00	43.103	5055.4	24.074	45.773	4556.0	21.700
215.00	43.843	5272.8	24.525	46.796	4788.3	22.271
220.00	44.585	5493.9	24.972	47.812	5024.9	22.840
225.00	45.331	5718.7	25.416	48.823	5266.5	23.406
230.00	46.085	5947.2	25.857	49.827	5513.1	23.970
235.00	46.852	6179.5	26.296	50.827	5764.7	24.531
240.00	47.635	6415.7	26.732	51.821	6021.3	25.089
245.00	48.429	6655.9	27.167	52.812	6282.9	25.645
250.00	49.226	6900.0	27.600	53.798	6549.4	26.198
255.00	50.019	7148.2	28.032	54.781	6820.9	26.749
260.00	50.804	7400.2	28.462	55.759	7097.2	27.297
265.00	51.586	7656.2	28.891	56.735	7378.5	27.843
270.00	52.371	7916.1	29.319	57.706	7664.6	28.387
275.00	52.870	8081.8	29.588	58.317	7847.3	28.729
280.00	53.166	8179.9	29.745	58.674	7955.5	28.929
285.00	53.981	8447.8	30.171	59.640	8251.3	29.469
290.00	54.223	8719.8	30.596	60.602	8551.9	30.007
295.00	55.693	8996.1	31.021	61.563	8857.3	30.543
298.15	57.156	9455.9	31.715	63.127	9365.5	31.412
300.00	57.488	9561.9	31.873	63.482	9482.6	31.609

H_0^0 IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Cole, A. G., Hutchens, J. O. and Stout, J. W.,
 Heat capacities from 11 to 305 K. and entropies of l-arginine.HCl,
 l-histidine.HCl, and l-lysine.HCl,
J. Phys. Chem. **67**, 2245-2247 (1963).

TABLE 14

MOLAL THERMODYNAMIC FUNCTIONS FOR L-ARGININE HYDROCHLORIDE
 $(\text{NH}_2\text{C}(\text{-NH}_2\text{Cl})\text{NH}(\text{CH}_2)_3\text{(NH}_2\text{)}\text{CHCOOH})$
 SOLID PHASE

GRAM MOLECULAR WT. = 210.66505 GRAMS
 $T \text{ DEG K} = 273.15 + T \text{ DEG C}$

1 CAL = 4.1840 JOULES

T DEG K	C_p^{C} CAL/DEG	$(H_T^0 - H_0^{\text{C}})$ CAL	$(H_T^0 - H_0^{\text{C}})/T$ CAL/DEG	S_T^0 CAL/DEG	$-(\alpha_T^0 - \alpha_0^{\text{C}})$ CAL	$-(G_T^0 - G_0^{\text{C}})/T$ CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.067	0.084	0.017	0.022	0.028	0.006
10.00	0.520	1.321	0.132	0.176	0.444	0.044
15.00	1.543	6.265	0.418	0.563	2.187	0.146
20.00	2.934	17.345	0.867	1.191	6.484	0.324
25.00	4.594	36.060	1.442	2.020	14.434	0.577
30.00	6.436	63.587	2.120	3.018	26.966	0.899
35.00	8.550	100.54	2.873	4.154	44.846	1.281
40.00	10.246	147.05	3.676	5.393	68.677	1.717
45.00	12.106	202.94	4.510	6.708	98.902	2.198
50.00	13.903	268.01	5.360	8.077	135.84	2.717
55.00	15.599	341.80	6.215	9.482	179.73	3.268
60.00	17.214	423.86	7.064	10.909	230.70	3.845
65.00	17.759	515.83	7.905	12.349	286.85	4.444
70.00	20.179	611.25	8.732	13.792	354.20	5.060
75.00	21.482	715.42	9.559	15.229	426.76	5.690
80.00	21.783	826.12	10.327	16.658	506.48	6.331
85.00	24.102	943.41	11.099	18.075	593.32	6.980
90.00	25.273	1066.9	11.854	19.420	687.25	7.636
95.00	26.254	1196.0	12.680	20.886	788.20	8.297
100.00	27.285	1330.4	13.304	22.264	896.08	8.961
105.00	28.436	1465.9	13.999	23.626	1010.8	9.627
110.00	29.466	1614.7	14.679	24.973	1132.3	10.294
115.00	30.496	1764.5	15.344	26.305	1260.5	10.961
120.00	31.411	1915.2	15.993	27.621	1395.3	11.628
125.00	32.303	2078.6	16.629	28.923	1536.7	12.294
130.00	33.306	2242.8	17.252	30.210	1684.5	12.958
135.00	34.266	2411.7	17.864	31.485	1838.8	13.621
140.00	35.174	2481.3	18.466	32.747	1999.4	14.281
145.00	36.174	2763.4	19.058	33.998	2166.2	14.940
150.00	36.486	2946.0	19.640	35.236	2339.3	15.596
155.00	37.845	3133.0	20.213	36.462	2518.6	16.249
160.00	39.731	3324.4	20.778	37.677	2703.9	16.900
165.00	39.616	3520.3	21.335	38.883	2895.3	17.547
170.00	40.498	3720.6	21.886	40.079	3092.7	18.193
175.00	41.368	3925.3	22.430	41.265	3296.1	18.835
180.00	42.224	4134.3	22.968	42.442	3505.4	19.474
185.00	43.073	4347.5	23.500	43.611	3720.5	20.111
190.00	43.923	4560.0	24.026	44.771	3941.5	20.745
195.00	44.775	4766.7	24.547	45.923	4168.2	21.375
200.00	45.624	5012.7	25.064	47.067	4400.7	22.003
205.00	46.463	5263.5	25.575	48.204	4638.9	22.629
210.00	47.314	5477.4	26.083	49.334	4882.7	23.251
215.00	48.161	5715.1	26.587	50.457	5132.2	23.871
220.00	49.012	5954.0	27.087	51.574	5387.3	24.488
225.00	49.867	6206.4	27.583	52.685	5647.9	25.102
230.00	50.714	6457.7	28.077	53.791	5914.1	25.714
235.00	51.568	6715.4	28.568	54.890	6185.8	26.323
240.00	52.413	6974.4	29.056	55.985	6463.0	26.929
245.00	53.261	7237.6	29.541	57.074	6745.7	27.533
250.00	54.117	7506.0	30.024	58.159	7033.8	28.135
255.00	54.963	7778.7	30.505	59.239	7327.2	28.734
260.00	55.817	8055.8	30.984	60.315	7626.1	29.331
265.00	56.713	8337.3	31.462	61.388	7930.4	29.926
270.00	57.604	8623.2	31.938	62.456	8240.0	30.519
275.00	58.151	8805.5	32.237	63.128	8437.8	30.891
275.00	58.470	8813.4	32.412	63.521	8555.0	31.109
280.00	59.26	9207.8	32.885	64.582	8875.2	31.697
285.00	60.176	9506.6	33.357	65.640	9200.8	32.283
290.00	61.022	9803.6	33.826	66.694	9531.6	32.868
295.00	61.864	10117.	34.294	67.744	9867.7	33.450
300.00	62.723	10313.	34.588	68.404	10082.	33.816
300.00	62.702	10428.	34.761	68.791	10209.	34.030

H_0^{C} IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Cole, A. G., Hutchens, J. O. and Stout, J. W.,
 Heat capacities from 11 to 305 K. and entropies of L-arginine.HCl,
 L-histidine.HCl, and L-lysine.HCl,
 J. Phys. Chem. 67, 2245-2247 (1963).

TABLE 15

MOLAL THERMODYNAMIC FUNCTIONS FOR L-HISTIDINE HYDROCHLORIDE
 $(C_9H_{13}N_2)(HCl)CH_2(NH_2)CHCOOH$
SOLID PHASE

GRAM MOLECULAR WT. = 191.61850 GRAMS
 $T \text{ DEG K} = 273.15 + T \text{ DEG C}$

1 CAL = 4.1840 JOULES

T DEG K	C_p^c CAL/DEG	$(H_T^0 - H_0^c)$ CAL	$(H_T^0 - H_0^c)/T$ CAL/DEG	S_T^0 CAL/DEG	$-(G_T^0 - G_0^c)$ CAL	$-(G_T^0 - G_0^c)/T$ CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.052	0.066	0.013	0.017	0.022	0.004
10.00	0.417	1.046	0.105	0.139	0.349	0.035
15.00	1.359	5.214	0.348	0.465	1.757	0.117
20.00	2.853	15.584	0.779	1.051	5.438	0.272
25.00	4.602	34.140	1.366	1.872	12.660	0.506
30.00	6.526	61.913	2.064	2.879	24.468	0.816
35.00	8.464	99.409	2.840	4.032	41.695	1.191
40.00	10.347	146.46	3.662	5.285	64.952	1.624
45.00	12.163	202.76	4.506	6.610	94.666	2.104
50.00	13.891	267.95	5.359	7.982	131.13	2.623
55.00	15.490	341.45	6.208	9.381	174.53	3.173
60.00	17.016	422.74	7.046	10.795	224.97	3.749
65.00	18.447	511.47	7.869	12.215	282.49	4.346
70.00	19.779	607.11	8.673	13.632	347.11	4.959
75.00	21.022	704.14	9.455	15.039	418.79	5.584
80.00	22.257	817.34	10.217	16.435	497.48	6.218
85.00	23.464	931.67	10.961	17.821	583.13	6.860
90.00	24.566	1051.8	11.687	19.194	675.67	7.507
95.00	25.563	1177.1	12.391	20.549	775.04	8.158
100.00	26.530	1307.4	13.074	21.885	881.13	8.811
105.00	27.490	1442.4	13.737	23.203	993.86	9.465
110.00	28.440	1582.3	14.384	24.504	1113.1	10.119
115.00	29.359	1726.8	15.015	25.788	1238.9	10.773
120.00	30.239	1875.8	15.632	27.056	1371.0	11.425
125.00	31.112	2024.2	16.233	28.309	1509.4	12.075
130.00	31.998	2186.9	16.823	29.546	1654.0	12.723
135.00	32.868	2349.1	17.401	30.770	1804.8	13.369
140.00	33.752	2515.6	17.968	31.981	1961.7	14.012
145.00	34.522	2686.1	18.525	33.178	2124.6	14.653
150.00	35.344	2860.8	19.072	34.362	2293.5	15.290
155.00	36.170	3039.6	19.610	35.534	2468.2	15.924
160.00	36.997	3222.5	20.140	36.695	2648.8	16.555
165.00	37.830	3409.5	20.664	37.847	2835.2	17.183
170.00	38.663	3600.8	21.181	38.986	3027.3	17.807
175.00	39.478	3796.1	21.692	40.121	3225.0	18.429
180.00	40.272	3995.5	22.197	41.244	3428.4	19.047
185.00	41.056	4198.8	22.696	42.358	3637.5	19.662
190.00	41.849	4406.1	23.190	43.464	3852.0	20.274
195.00	42.656	4617.4	23.679	44.561	4072.1	20.882
200.00	43.473	4832.7	24.163	45.651	4297.6	21.488
205.00	44.293	5052.1	24.644	46.735	4528.6	22.091
210.00	45.112	5275.6	25.122	47.812	4765.0	22.690
215.00	45.924	5503.2	25.596	48.883	5006.7	23.287
220.00	46.726	5734.8	26.067	49.946	5253.8	23.881
225.00	47.517	5970.4	26.535	51.007	5506.2	24.472
230.00	48.301	6210.0	27.000	52.060	5763.8	25.060
235.00	49.089	6453.5	27.462	53.107	6026.8	25.646
240.00	49.891	6700.9	27.920	54.149	6294.9	26.229
245.00	50.713	6952.4	28.377	55.186	6568.2	26.809
250.00	51.558	7208.1	28.832	56.219	6846.8	27.387
255.00	52.417	7468.0	29.286	57.249	7130.4	27.962
260.00	53.277	7732.2	29.739	58.275	7419.2	28.536
265.00	54.127	8000.8	30.192	59.298	7713.2	29.106
270.00	54.961	8273.5	30.643	60.317	8012.2	29.675
273.15	55.478	8447.4	30.926	60.958	8203.2	30.032
275.00	55.780	8550.3	31.092	61.333	8316.3	30.241
280.00	56.191	8681.3	31.540	62.346	8625.5	30.805
285.00	57.399	9116.2	31.987	63.355	8939.8	31.368
290.00	58.210	9405.3	32.432	64.360	9259.1	31.928
295.00	59.028	9698.4	32.876	65.362	9583.4	32.486
298.15	59.549	9885.1	33.155	65.992	9790.3	32.837
300.00	59.858	9995.6	33.319	66.361	9912.7	33.042

 H_0^c IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Cole, A. G., Hutchens, J. O. and Stout, J. W.,
Heat capacities from 11 to 305 K. and entropies of L-arginine.HCl,
L-histidine.HCl, and L-lysine.HCl,
J. Phys. Chem. 67, 2245-2247 (1963).

TABLE 16

MOLAL THERMODYNAMIC FUNCTIONS FOR L-CYSTINE
 $((\text{HOOC}(\text{NH}_2)\text{CHCH}_2\text{S}-)_2$
 SOLID PHASE

GRAM MOLECULAR WT. = 240.30154 GRAMS
 T DEG K = 273.15 + T DEG C

1 CAL = 4.1840 JOULES

T DEG K	c_p^C CAL/DEG	$(H_T^0 - H_0^C)$ CAL	$(H_T^0 - H_0^C)/T$ CAL/DEG	s_T^0 CAL/DEG	$-(G_T^0 - H_0^C)$ CAL	$-(G_T^0 - H_0^C)/T$ CAL/DEG
0.00	0.000	0.000	0.000	0.000	0.000	0.000
5.00	0.050	0.063	0.013	0.017	0.021	0.004
10.00	0.103	0.092	0.029	0.032	0.033	0.003
15.00	0.164	0.088	0.026	0.043	0.062	0.011
20.00	0.260	0.109	0.025	0.081	0.104	0.025
25.00	0.311	0.133	0.027	0.147	0.180	0.047
30.00	0.395	0.181	0.027	0.269	0.286	0.076
35.00	0.419	0.202	0.033	0.376	0.398	0.113
40.00	0.701	0.204	0.043	0.451	0.691	0.151
45.00	1.142	0.197	0.0217	0.183	0.477	0.096
50.00	1.313	0.213	0.023	0.475	1.246	0.245
55.00	1.469	0.209	0.031	0.799	1.632	0.296
60.00	1.623	0.202	0.034	1.044	2.106	0.511
65.00	1.773	0.206	0.040	1.150	2.647	0.673
70.00	1.912	0.188	0.0217	1.286	3.267	0.652
75.00	2.044	0.170	0.0188	1.424	3.934	0.246
80.00	21.783	0.166	0.0176	1.559	4.680	0.850
85.00	24.096	0.188	0.0193	1.695	5.493	0.463
90.00	24.305	0.104	0.0127	1.831	6.375	0.084
95.00	25.427	0.134	0.0194	1.965	7.324	0.710
100.00	26.515	0.126	0.0164	2.098	8.340	0.341
105.00	27.600	0.139	0.0133	2.230	9.423	0.975
110.00	28.681	0.150	0.0106	2.361	10.572	0.610
115.00	29.737	0.167	0.0167	2.491	11.785	0.248
120.00	30.758	0.187	0.0131	2.620	13.063	0.886
125.00	31.769	0.194	0.0195	2.747	14.405	0.524
130.00	32.786	0.215	0.0158	2.874	15.810	1.162
135.00	33.796	0.232	0.0201	3.000	17.279	0.799
140.00	34.763	0.245	0.0181	3.124	18.810	1.436
145.00	35.740	0.269	0.0143	3.248	20.404	0.071
150.00	35.694	0.261	0.0106	3.371	22.059	1.706
155.00	37.653	0.306	0.0292	3.493	23.775	0.338
160.00	38.611	0.327	0.0172	3.614	25.552	0.970
165.00	39.660	0.342	0.0175	3.734	27.389	0.599
170.00	40.647	0.323	0.0131	3.853	29.286	0.227
175.00	41.425	0.387	0.0174	3.972	31.242	0.853
180.00	42.359	0.407	0.0243	4.096	33.258	0.477
185.00	42.275	0.421	0.0296	4.207	35.333	0.099
190.00	44.179	0.470	0.0252	4.324	37.466	0.719
195.00	45.072	0.493	0.0207	4.440	39.657	0.317
200.00	46.052	0.490	0.0260	4.555	41.906	0.953
205.00	46.822	0.512	0.0135	4.670	44.213	0.567
210.00	47.693	0.538	0.0262	4.784	46.577	0.179
215.00	48.567	0.562	0.0184	4.897	48.997	0.789
220.00	49.439	0.584	0.0203	5.0100	51.474	0.397
225.00	50.304	0.612	0.0218	5.1221	54.007	0.003
230.00	51.162	0.637	0.0279	5.2336	56.596	0.607
235.00	52.011	0.663	0.0236	5.3445	59.240	0.209
240.00	52.581	0.687	0.0287	5.4549	61.940	0.808
245.00	53.749	0.716	0.0242	5.5649	64.695	0.406
250.00	54.612	0.745	0.0274	5.6743	67.505	0.002
255.00	55.487	0.771	0.0237	5.7833	70.369	0.596
260.00	56.324	0.799	0.0271	5.8919	73.288	0.188
265.00	57.166	0.823	0.0222	6.0000	76.261	0.778
270.00	57.992	0.856	0.0170	6.1076	79.288	0.366
275.00	58.509	0.875	0.0216	6.1752	81.223	0.736
280.00	58.813	0.883	0.0195	6.2148	82.369	0.952
285.00	59.633	0.914	0.0267	6.3215	85.503	0.537
290.00	60.455	0.945	0.0215	6.4277	88.690	0.119
295.00	61.280	0.975	0.0236	6.5336	91.930	0.700
298.15	62.105	1.006	0.0111	6.6391	95.224	0.279
300.00	62.622	1.025	0.0410	6.7053	97.325	0.643
300.00	62.974	1.037	0.0585	6.7441	98.569	0.856

H_0^C IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Huffman, H. M. and Ellis, E. L.,
 Thermal data. III. The heat capacities, entropies and free energies of
 four organic compounds containing sulfur,
 J. Am. Chem. Soc. 57, 46-48 (1935).

Hutchens, J. O., Cole, A. G. and Stout, J. W.,
 Heat capacities and entropies of l-cystine and l-methionine,
 J. Biol. Chem. 239, 591-595 (1964).

TABLE 17							
MOLAL THERMODYNAMIC FUNCTIONS FOR L-PROLINE							
(C ₄ H ₈ NCOOH)							
SOLID PHASE							
GRAM MOLECULAR WT. = 115.13298 GRAMS				1 CAL = 4.1840 JOULES			
T DEG K = 273.15 + T DEG C				1 CAL = 4.1840 JOULES			
T	c _p ^C	(H _T ⁰ -H ₀ ^C)	(H _T ⁰ -H ₀ ^C)/T	S _T ⁰	-(G _T ⁰ -H ₀ ^C)	-(G _T ⁰ -H ₀ ^C)/T	
DEG K	CAL/DEG	CAL	CAL/DEG	CAL/DEG	CAL	CAL/DEG	
0.00	0.000	0.000	0.000	0.000	0.000	0.000	
5.00	0.039	0.049	0.010	0.013	0.016	0.003	
10.00	0.310	0.780	0.078	0.104	0.261	0.026	
15.00	0.981	3.828	0.255	0.342	1.305	0.087	
20.00	1.989	11.161	0.558	0.757	3.982	0.199	
25.00	3.114	23.900	0.956	1.321	9.126	0.365	
30.00	4.267	42.349	1.412	1.991	17.368	0.579	
35.00	5.411	66.560	1.902	2.735	29.155	0.833	
40.00	6.494	96.553	2.409	3.529	44.798	1.120	
45.00	7.512	141.39	2.920	4.353	64.493	1.433	
50.00	8.459	171.56	3.427	5.194	88.356	1.767	
55.00	9.320	215.83	3.924	6.041	118.44	2.117	
60.00	10.144	264.51	4.408	6.888	148.77	2.479	
65.00	10.912	317.17	4.880	7.731	185.32	2.851	
70.00	11.607	373.51	5.336	8.565	226.06	3.229	
75.00	12.272	433.21	5.776	9.389	270.95	3.613	
80.00	12.934	495.24	6.203	10.202	317.93	3.999	
85.00	13.553	562.48	6.617	11.005	372.96	4.388	
90.00	14.129	631.64	7.019	11.746	424.46	4.777	
95.00	14.675	703.72	7.408	12.570	490.40	5.167	
100.00	15.194	778.40	7.784	13.341	555.69	5.557	
105.00	15.703	855.64	8.149	14.095	624.29	5.946	
110.00	16.215	939.43	8.504	14.837	696.62	6.333	
115.00	16.731	1017.8	8.850	15.569	772.64	6.719	
120.00	17.244	1102.7	9.189	16.292	852.29	7.102	
125.00	17.748	1190.2	9.522	17.006	935.54	7.484	
130.00	18.256	1280.4	9.848	17.712	1022.3	7.864	
135.00	18.767	1372.6	10.169	18.411	1112.6	8.242	
140.00	19.275	1467.4	10.485	19.102	1206.4	8.617	
145.00	19.787	1565.5	10.797	19.788	1303.7	8.991	
150.00	20.314	1665.8	11.105	20.467	1404.3	9.362	
155.00	20.844	1768.7	11.411	21.142	1504.3	9.731	
160.00	21.374	1874.3	11.714	21.412	1612.7	10.098	
165.00	21.907	1982.5	12.015	22.478	1720.4	10.465	
170.00	22.438	2093.3	12.314	23.140	1840.5	10.826	
175.00	22.976	2206.9	12.611	23.798	1957.8	11.188	
180.00	23.518	2323.1	12.906	24.453	2076.5	11.547	
185.00	24.050	2442.0	13.200	25.109	2202.4	11.905	
190.00	24.597	2560.7	13.493	25.754	2329.5	12.261	
195.00	25.142	2688.0	13.785	26.499	2459.9	12.615	
200.00	25.641	2814.9	14.075	27.042	2593.5	12.968	
205.00	26.156	2944.4	14.363	27.682	2730.3	13.319	
210.00	26.678	3076.5	14.650	28.318	2870.3	13.665	
215.00	27.213	3211.2	14.936	28.952	3013.5	14.016	
220.00	27.745	3349.6	15.221	29.584	3153.8	14.363	
225.00	28.274	3486.6	15.505	30.213	3299.3	14.728	
230.00	28.793	3631.3	15.786	30.840	3462.0	15.052	
235.00	29.306	3776.6	16.070	31.468	3617.7	15.495	
240.00	29.824	3924.4	16.352	32.087	3776.6	15.736	
245.00	30.340	4074.8	16.632	32.708	3938.6	16.076	
250.00	30.857	4221.9	16.912	33.326	4103.7	16.415	
255.00	31.425	4365.7	17.191	33.943	4271.9	16.752	
260.00	31.969	4514.2	17.470	34.559	4443.1	17.089	
265.00	32.505	4705.4	17.749	35.173	4617.4	17.424	
270.00	33.037	4867.4	18.027	35.785	4794.8	17.759	
275.00	33.571	5027.1	18.202	36.171	4908.2	18.099	
280.00	33.968	5032.7	18.304	36.396	4975.3	18.092	
285.00	34.412	5102.9	18.482	37.006	5158.8	18.424	
290.00	34.842	5185.9	18.859	37.614	5345.3	18.756	
295.00	35.240	5247.3	19.156	38.222	5534.9	19.086	
300.00	35.646	5266.7	19.412	38.828	5721.6	19.415	
305.00	36.046	53839.8	19.587	39.410	5850.5	19.623	
310.00	36.319	5906.8	19.689	39.434	5923.2	19.744	

^C IS THE ENTHALPY OF THE SOLID AT 0 DEG K AND 1 ATM PRESSURE.

Huffman, H. M. and Fox, S. W.,
Thermal data. XIII. The heat capacities and entropies of creatine hydrate,
di-citrulline, di-ornithine, L-proline and taurine,
J. Am. Chem. Soc. 62, 3464-3465 (1940).

Cole, A. G., Hutchens, J. O. and Stout, J. W.,
Heat capacities from 11 to 305°K. and entropies of L-phenylalanine,
L-proline, L-tryptophane, and L-tyrosine. Some free energies of formation,
J. Phys. Chem. 67, 1852-1855 (1963).

Section II

Heats and Free Energies of Formation of Compounds of C, H, N, O, P, and S

E. S. Domalski and George T. Armstrong

The data on the accompanying tables were obtained by a search of the references listed, each of which is a competent review of thermodynamic data covering many of the compounds of interest. Where data were available in Reference 1 they were used. Values found in Reference 2 were taken if information was not found in the other references. The list is given in the Appendix of NBS Report 8521 and was used as a basis for the search. Only a few compounds were included in the table which were not on the original list. On this account, the list of compounds is by no means complete, and it will be augmented in the future. While data in Table 1 may be expected to be the best available for most of the compounds, new data may be available for a few, and not have been included in the reviews searched. Estimates of the uncertainties which should be ascribed to the data have not yet been made here. Absence of data for a compound listed in the table does not necessarily, at this stage of the study, mean that no measurements have been made on the compound. We have had no way of indicating partial data, insufficient for calculation of enthalpy or free energy of formation. In addition, as mentioned before, very recent publications have not been covered thus far in the search.

TABLE I
Preliminary Table of Selected Thermodynamic Data on Compounds of CHNOPS
Containing Not More Than One C Atom per Molecule

Empirical Formula	Functional Group Formula	Name	State	ΔH_{f298}°	ΔF_{f298}°	S_{298}°	Ref.
				kcal mole ⁻¹	kcal mole ⁻¹	cal mole ⁻¹ deg ⁻¹	
C	C	carbon, monatomic	g	171.291	160.442	37.760	[1]
		graphite	c	0	0	1.372	[1]
		diamond	c	0.4533	0.6930	0.568	[1]
CH	CH	methylidyne	g	142.1			[4]
CHN	HCN	hydrogen cyanide	g	31.2	28.7	48.23	[2]
		hydrocyanic acid,	m=l	25.2	29.0	26.97	[2]
CHNO	HCNO	hydrogen cyanate	g				
		cyanic acid,	m=l	aq	-35.1	-28.9	43.6
CHNS	HCNS	hydrogen thiocyanate	m=l	g			
		thiocyanic acid,	m=l	aq	17.7		
CH ₃ O ₆	CH(NO ₂) ₃	trinitromethane	l	-18.6			[2]
CHO	CHO	formyl	g	-2,900	-6.543	53.683	[4]
CH ₂	CH ₂	methylene	g	95.000	91.809	43.271	[11]
CH ₂ N ₂	(NH ₂)CN	cyanamide	800 H ₂ O	c	9.2		[2]
				aq	12.9		[2]
CH ₂ N ₂	(CH ₂)N ₂	diazomethane	g				
CH ₂ N ₂ O ₃	(NO ₂)CH(NOH)	formonitrolic acid	c				
CH ₂ N ₂ O ₄	CH ₂ (NO ₂) ₂	dinitromethane	l				
CH ₂ N ₄	NHN:NCH:N	1,2,3,5-tetrazole	c				
CH ₂ O	HCHO	formaldehyde	60 H ₂ O	g	-27.700	-26.266	52.261
			40 CH ₃ OH	aq	-42.5		[4]
				aq	-42.7		[2]
(CH ₂ O) _x	(CH ₂ O) _x	paraformaldehyde	c				
CH ₂ O ₂	HCOOH	formic acid	g	-86.67	-80.24	60.0	[2]
			l	-97.8	-82.7	30.82	[2]
			aq	-98.0	-85.1	39.1	[2]
			0.2 H ₂ O	aq	-97.86		[2]
			0.5 H ₂ O	aq	-97.93		[2]
			1.0 H ₂ O	aq	-98.00		[2]
			1.5 H ₂ O	aq	-98.01		[2]
			2.0 H ₂ O	aq	-97.99		[2]
			2.5 H ₂ O	aq	-97.99		[2]
			4 H ₂ O	aq	-97.98		[2]
CH ₂ O ₃	H ₂ CO ₃	carbonic acid undissociated, m=l	aq	-97.96			
			0.2 H ₂ O	aq	-97.94		[2]
			0.5 H ₂ O	aq	-97.93		[2]
			1.0 H ₂ O	aq	-97.93		[2]
			1.5 H ₂ O	aq	-97.93		[2]
			2.0 H ₂ O	aq	-97.94		[2]
			2.5 H ₂ O	aq	-97.94		[2]
			50 H ₂ O	aq	-97.95		[2]
			100 H ₂ O	aq	-97.96		[2]
			200 H ₂ O	aq	-97.97		[2]
CH ₂ S ₃	H ₂ CS ₃	trithiocarbonic acid	aq	-98.0			
CH ₃	CH ₃	methyl	g	31.940	32.546	46.137	[8]
CH ₃ NO	HCONH ₂	formamide	l	-61.6			[2]
			aq	-56.0			[2]
CH ₃ NO	H ₂ C(NOH)	formaldehyde oxime	l				
CH ₃ NO ₂	CH ₃ NO ₂	nitromethane	l	-21.28	2.26	41.1	[2]
			aq	-20.7			[2]

Table I. Selected Thermodynamic Data (Cont.)

Empirical Formula	Functional Group Formula	Name	State	ΔH_f° kcal mole ⁻¹	ΔF_f° kcal mole ⁻¹	S_f° cal mole ⁻¹ deg ⁻¹	Ref.	
CH ₅ NO	CH ₃ ONH ₂	methoxyamine	c aq					
CH ₅ NO ₂	HCOONH ₄	ammonium formate	c aq	-132.8 -129.83			[2] [2]	
CH ₅ NO ₃	NH ₄ HCO ₃	ammonium bicarbonate	m=l	c aq	-203.7 -196.92	-159.31	49.7	[2] [2]
CH ₅ N ₃	NH ₂ C(NH)NH ₂	guanidine		c aq	-17.0 -18.3			[2] [2]
CH ₅ N ₃ O	NH ₂ CONHNH ₂	semicarbazide	c aq					
CH ₅ N ₃ O ₃ S	NH ₂ CSNH ₂ HNO ₃	thiourea nitrate	c	-74.5			[2]	
CH ₅ N ₃ O ₄	NH ₂ CONH ₂ HNO ₃	urea nitrate	c aq	-114.8			[2]	
CH ₅ N ₃ S	NH ₂ CSNHNH ₂	thiosemicarbazide	c aq					
CH ₅ O ₃ P	CH ₃ PO(OH) ₂	methyl phosphonic acid	c aq					
CH ₅ P	CH ₃ PH ₂	methyl phosphine	g					
CH ₆ N ₂	CH ₃ NHNH ₂	methyl hydrazine	g					
CH ₆ N ₂ O ₂	NH ₂ COONH ₄	ammonium carbamate	c aq	-154.21 -150.4	-109.47	39.70	[2] [2]	
CH ₆ N ₄	NH ₂ C(NH)NHNH ₂	1-aminoguanidine	c aq					
CH ₆ N ₄ O	CO(NHNH ₂) ₂	carbohydrazide	c					
CH ₈ N ₂ O ₃	(NH ₄) ₂ CO ₃	ammonium carbonate	c aq	-225.11	-164.22	41.2	[2]	
CH ₁₂ O ₈	CO ₂ •6H ₂ O	carbon dioxide hexahydrate	c	-520			[2]	
CH ₁₆ O ₆	CH ₄ •6H ₂ O	methane hexahydrate	c	-445			[2]	
CN	CN	cyano	g	109.000	101.796	48.406	[8]	
(CN) _x	(CN) _x	paracyanogen	c					
CN ₄	CN(N ₃)	cyanogen azide	c	92.6			[2]	
$\frac{1}{x}$ (CN ₄) _x	$\frac{1}{x}$ (CN(N ₃)) _x	paracyanogen azide	c	82.2			[2]	
CN ₄ O ₈	C(NO ₂) ₄	tetranitromethane	l	8.8			[2]	
CO	CO	carbon monoxide	g	-26.416	-32.780	42.214	[1]	
COS	COS	carbonyl sulfide	g	-33.080	-39.589	55.323	[4]	
CO ₂	CO ₂	carbon dioxide	g	-94.051	-94.261	51.072	[1]	
		undissociated, m=l	aq	-98.85	-92.26	28.3	[1]	
CP	CP	carbon phosphide	g	111.700	98.327	51.661	[6]	
CS	CS	carbon monosulfide	g	55.000	42.684	50.299	[8]	
CS ₂	CS ₂	carbon disulfide	l	27.980	15.991	56.832	[5]	
C ₂	C ₂	carbon diatomic	g	199.026	185.636	47.628	[6]	
C ₂ H ₈ N ₂ O ₄	N ₂ H ₄ •(HCOOH) ₂	hydrazine formate	c aq					
C ₂ N ₂	(CN) ₂	cyanogen	g	73.870	71.117	57.711	[4]	

Table I. Selected Thermodynamic Data (Cont.)

Empirical Formula	Functional Group Formula	Name	State	ΔH_f°	ΔF_f°	S_f°	Ref.
				kcal mole ⁻¹	kcal mole ⁻¹	cal mole ⁻¹ deg ⁻¹	
C ₃	C ₃	carbon, triatomic	g	189.670	175.777	50.688	[3]
C ₃ O ₂	C ₃ O ₂	carbon suboxide	g	-8.300	-10.726	61.236	[3]
C ₃ S ₂	C ₃ S ₂	carbon subsulfide	g				
C ₄	C ₄	carbon, tetratomic	g	242.321	226.629	58.083	[3]
C ₄ N ₂	C ₂ (CN) ₂	carbon subnitride	g	122.900	113.575	67.936	[3]
C ₅	C ₅	carbon, pentatomic	g	242.374	226.634	59.608	[3]
H	H	hydrogen monatomic	g	52.095	48.580	27.391	[1]
HN	HN	imideogen	g	79.200	77.765	43.297	[3]
HN ₃	HN ₃	hydrogen azide	g	70.3	78.4	57.09	[1]
		hydrazoic acid undissociated, m=1	aq	63.1 62.16	78.2 76.9	33.6 34.9	[1]
HNO	HNO	nitroxyl	g	23.800	26.859	52.729	[9]
HNO ₂	HNO ₂	cis hydrogen nitrite trans hydrogen nitrite hydrogen nitrite (cis-trans mixture)	g	-18.64 -19.15	-10.27 -10.82	59.43 59.54	[1]
		nitrous acid	aq	-19.0 -28.5	-11.0 -13.3	60.7 36.5	[1]
HNO ₃	HNO ₃	hydrogen nitrate	g	-32.28	-17.87	63.64	[1]
		nitric acid, m=1	l	-41.61	-19.31	37.19	[1]
			aq	-49.56	-26.61	35.0	[1]
		1 H ₂ O	aq	-44.845			
		2 H ₂ O	aq	-46.500			
		3 H ₂ O	aq	-47.459			
		5 H ₂ O	aq	-48.462			
		10 H ₂ O	aq	-49.192			
		25 H ₂ O	aq	-49.430			
		50 H ₂ O	aq	-49.439			
		100 H ₂ O	aq	-49.440			
		500 H ₂ O	aq	-49.468			
		1000 H ₂ O	aq	-49.484			
		5000 H ₂ O	aq	-49.518			
		10000 H ₂ O	aq	-49.529			
		50000 H ₂ O	aq	-49.545			
H ₃ NO ₄	HNO ₃ ·H ₂ O	nitric acid hydrate	t	-112.960	-78.410	51.83	[2]
HNO ₅ S	(NO)HSO ₄	nitrosyl sulfuric acid	c				
HO	OH	hydroxyl	g	9.31	8.18	43.890	[1]
HO ₂	HO ₂	hydroperoxy	g	5.000	8.049	54.383	[10]
HP	PH	phosphorus monohydride	g	59.170	51.467	46.891	[7]
HP ₂	P ₂ H	diphosphorus monohydride	c	-14.5			[2]
HPO ₃	HPO ₃	metaphosphoric acid	aq	-226.7 -233.5			[1]
HS	SH	sulfur monohydride	g	32.000	24.990	46.745	[3]
H ₂		hydrogen	g	0 -1.0	0 4.2	31.208 15.8	[1]
H ₂ N	NH ₂	amidogen	g	40.300	42.976	45.113	[3]
H ₂ N ₂ O ₂	H ₂ N ₂ O ₂	hyponitrous acid, m=1	aq	-13.7	8.4	52	[2]
H ₂ O	H ₂ O	water	g	-57.796 -68.315	-54.635 -56.688	45.104 16.71	[1]
H ₂ O ₂	H ₂ O ₂	hydrogen peroxide	g	-32.58	-25.25	55.6	[1]
		m=1	aq	-44.88	-28.78	26.2	[1]
		0.5 H ₂ O	aq	-45.69	-32.05	34.4	[1]
		1 H ₂ O	aq	-45.198			
		5 H ₂ O	aq	-45.365			
		10 H ₂ O	aq	-45.638			
		50 H ₂ O	aq	-45.670			
			aq	-45.687			

Table I. Selected Thermodynamic Data (Cont.)

Empirical Formula	Functional Group Formula	Name	State	ΔH_f° 298 kcal mole ⁻¹	ΔF_f° 298 kcal mole ⁻¹	S° 298 cal mole ⁻¹ deg ⁻¹	Ref.
H ₂ P	PH ₂	phosphorus dihydride	g	30.100	25.884	50.800	[12]
H ₂ S	H ₂ S	hydrogen sulfide hydrogensulfuric acid	g aq	-4.93 -9.5	-8.02 -6.66	49.16 29	[1] [1]
H ₂ S ₂	H ₂ S ₂	hydrogen disulfide	l	-5.5			[2]
H ₂ S ₃	H ₂ S ₃	hydrogen trisulfide	g				
H ₂ S ₅	H ₂ S ₅	hydrogen pentasulfide	l	0.7			[2]
H ₂ SO ₃	H ₂ SO ₃	sulfurous acid undissociated, m=1	aq	-145.51	-128.56	55.5	[1]
		100 H ₂ O	aq	-146.369			[1]
		200 H ₂ O	aq	-146.670			[1]
		500 H ₂ O	aq	-147.126			[1]
		1000 H ₂ O	aq	-147.516			[1]
		2000 H ₂ O	aq	-147.957			[1]
		5000 H ₂ O	aq	-148.524			[1]
		10000 H ₂ O	aq	-148.899			[1]
H ₂ SO ₄	H ₂ SO ₄	sulfuric acid	l m=1 aq	-194.548 -217.32	-164.942 -177.97	37.501 4.8	[1] [1]
		1 H ₂ O	aq	-201.193			[1]
		2 H ₂ O	aq	-204.425			[1]
		3 H ₂ O	aq	-206.241			[1]
		4 H ₂ O	aq	-207.428			[1]
		5 H ₂ O	aq	-208.288			[1]
		6 H ₂ O	aq	-208.944			[1]
		8 H ₂ O	aq	-209.865			[1]
		10 H ₂ O	aq	-210.451			[1]
		15 H ₂ O	aq	-211.191			[1]
		25 H ₂ O	aq	-211.660			[1]
		50 H ₂ O	aq	-211.941			[1]
		75 H ₂ O	aq	-212.068			[1]
		100 H ₂ O	aq	-212.150			[1]
		115 H ₂ O	aq	-212.192			[1]
		200 H ₂ O	aq	-212.387			[1]
		300 H ₂ O	aq	-212.565			[1]
		500 H ₂ O	aq	-212.833			[1]
		800 H ₂ O	aq	-213.128			[1]
		1000 H ₂ O	aq	-213.275			[1]
		1500 H ₂ O	aq	-213.552			[1]
		2000 H ₂ O	aq	-213.740			[1]
		3000 H ₂ O	aq	-214.015			[1]
		5000 H ₂ O	aq	-214.390			[1]
		10000 H ₂ O	aq	-215.060			[1]
		20000 H ₂ O	aq	-215.880			[1]
		50000 H ₂ O	aq	-216.545			[1]
		100000 H ₂ O	aq	-216.875			[1]
		500000 H ₂ O	aq	-217.189			[1]
H ₂ SO ₅	H ₂ SO ₅	peroxymonosulfuric acid	c				
H ₂ S ₂ O ₄	H ₂ S ₂ O ₄	dithionous acid	aq	-164			[2]
H ₂ S ₂ O ₆	H ₂ S ₂ O ₆	dithionic acid	aq	-280.0			[2]
H ₂ S ₂ O ₇	H ₂ S ₂ O ₇	pyrosulfuric acid	c	-304.4			[1]
H ₂ S ₂ O ₈	H ₂ S ₂ O ₈	peroxydisulfuric acid, m=1	aq	-320.0	-265.4	59.3	[2]
H ₃ N	NH ₃	ammonia undissociated, m=1	g aq	-11.07 -19.19	-3.94 -6.35	45.97 26.6	[1] [1]
		1 H ₂ O	aq	-18.011			[1]
		2 H ₂ O	aq	-18.560			[1]
		5 H ₂ O	aq	-18.945			[1]
		10 H ₂ O	aq	-19.074			[1]
		20 H ₂ O	aq	-19.125			[1]
		50 H ₂ O	aq	-19.156			[1]
		100 H ₂ O	aq	-19.167			[1]
		500 H ₂ O	aq	-19.173			[1]
		1000 H ₂ O	aq	-19.171			[1]
		5000 H ₂ O	aq	-19.154			[1]
		10000 H ₂ O	aq	-19.140			[1]
		50000 H ₂ O	aq	-19.086			[1]

Table I. Selected Thermodynamic Data (Cont.)

Table I. Selected Thermodynamic Data (Cont.)

Empirical Formula	Functional Group Formula	Name	State	$\Delta H_f^{\circ} \text{298}$ kcal mole ⁻¹	$\Delta F_f^{\circ} \text{298}$ kcal mole ⁻¹	$S^{\circ} \text{298}$ cal mole ⁻¹ deg ⁻¹	Ref.
H ₅ NO	NH ₄ OH	ammonium hydroxide undissociated, m=1 ionized, m=1	l aq aq	-86.33 -87.505 -86.64	-60.74 -63.04 -56.56	39.57 43.3 24.5	[1] [1] [1]
H ₅ NS	NH ₄ SH	ammonium hydrosulfide 200 H ₂ O	c aq	-37.5 -34.8	-12.1	23.3	[1] [1]
H ₅ NO ₃ S	NH ₄ HSO ₃	ammonium bisulfite 300 H ₂ O	c aq	-183.7 -181.3			[1] [1]
H ₅ NO ₄ S	NH ₄ HSO ₄	ammonium bisulfate 200 H ₂ O	c aq	-245.45 -245.65			[1] [1]
H ₅ NO ₅ S	NH ₂ OH•H ₂ SO ₄	hydroxylamine sulfate	aq	-246.7			[1]
H ₅ N ₃ O ₃	N ₂ H ₄ •HNO ₃	hydrazine nitrate m=1	c aq	-60.13 -51.41	-6.91	71	[1] [1]
H ₆ NO ₂ P	(NH ₄) ₂ PO ₂	ammonium hypophosphite	c aq				
H ₆ NO ₃ P	(NH ₄) ₂ PO ₃	ammonium orthophosphite	c aq				
H ₆ NO ₄ P	(NH ₄) ₂ PO ₄	primary ammonium orthophosphate m=1 15 H ₂ O 50 H ₂ O 100 H ₂ O 500 H ₂ O 1000 H ₂ O ∞ H ₂ O	c aq aq aq aq aq aq	-345.94 -342.05 -342.157 -342.113 -342.088 -342.059 -342.055 -342.05	-289.89 -289.70	36.32 48.7	[1] [1] [1] [1] [1] [1] [1]
H ₆ N ₂ O	N ₂ H ₄ •H ₂ O	hydrazine hydrate undissociated, m=1	g l aq	-49.0 -58.01 -60.11	-18.9 -26.1	63 49.7	[1] [1] [1]
H ₆ N ₂ O ₃ S	(NH ₂)SO ₃ NH ₄	ammonium sulfamate	c aq				
H ₆ N ₂ O ₄ S	N ₂ H ₄ •H ₂ SO ₄	hydrazine sulfate 1000 H ₂ O	c aq	-231.6 -223.44			[2] [2]
H ₆ P ₁₂	(P ₂ H ₄) ₃	diphosphine trimer	c				
H ₆ •5O ₃ •5P ₂	H ₄ P ₂ O ₇ • ³ H ₂ O	pyrophosphoric acid hydrate	c l	-640.9 -637.3			[1] [1]
H ₇ NO ₆	HNO ₃ •3H ₂ O	nitric acid trihydrate	l	-252.203	-193.701	82.92	[2]
H ₇ N ₂ O ₃ P	N ₂ H ₄ •H ₃ PO ₃	hydrazine orthophosphite	c aq				
H ₇ N ₂ O ₄ P	N ₂ H ₄ •H ₃ PO ₄	hydrazine orthophosphate	c aq				
H ₈ N ₂ O ₃ S	(NH ₄) ₂ SO ₃	ammonium sulfite 400 H ₂ O	c aq	-211.6 -211.0			[1] [1]
H ₈ N ₂ O ₃ S ₂	(NH ₄) ₂ S ₂ O ₃	ammonium thiosulfate	c aq				
H ₈ N ₂ O ₄ S	(NH ₄) ₂ SO ₄	ammonium sulfate m=1 10 H ₂ O 50 H ₂ O 100 H ₂ O 500 H ₂ O 1000 H ₂ O	c aq aq aq aq aq	-282.23 -280.66 -280.72 -280.51 -280.407 -280.242 -280.217	-215.56 -215.77	52.6 58.6	[1] [1] [1] [1] [1] [1] [1]
H ₈ N ₂ O ₅ S ₂	(NH ₄) ₂ S ₂ O ₅	ammonium pyrosulfite, m=1	aq	-295.3			[2]
H ₈ N ₂ O ₆ P ₂	N ₂ H ₄ •H ₄ P ₂ O ₆	hydrazine hypophosphite	c aq				
H ₈ N ₂ O ₆ S	2NH ₂ OH•H ₂ SO ₄	dihydroxylamine sulfate	aq	-281.3			[1]
H ₈ N ₂ O ₆ S ₂	(NH ₄) ₂ S ₂ O ₆	ammonium dithionate	c aq				

Table I. Selected Thermodynamic Data (Cont.)

Empirical Formula	Functional Group Formula	Name	State	ΔH_f° kcal mole ⁻¹	ΔF_f° kcal mole ⁻¹	S_f° cal mole ⁻¹ deg ⁻¹	Ref.
H ₈ N ₂ O ₇ S ₂	(NH ₄) ₂ S ₂ O ₇	ammonium pyrosulfate	c aq				
H ₈ N ₂ O ₈ S ₂	(NH ₄) ₂ S ₂ O ₈	ammonium peroxydisulfate	c aq	-392.5 -383.3			[1] [1]
H ₈ N ₂ S	(NH ₄) ₂ S	ammonium monosulfide	aq	-54.5			[2]
H ₈ N ₂ S ₄	(NH ₄) ₂ S ₄	ammonium tetrasulfide	c aq	-67.4 -60.0			[2] [2]
H ₈ N ₂ S ₅	(NH ₄) ₂ S ₅	ammonium pentasulfide	c aq	-68.8 -61.2			[2] [2]
H ₈ O ₄ SP ₂	(PH ₄) ₂ SO ₄	phosphonium sulfate	c aq				
H ₉ N ₂ O ₄ P	(NH ₄) ₂ HPO ₄	secondary ammonium orthophosphate	c aq 15 H ₂ O 50 H ₂ O 100 H ₂ O 500 H ₂ O 1000 H ₂ O	-374.50 -372.71 -370.40 -370.85 -371.22 -371.4 -371.44	-298.85	46.2	[1] [1] [1] [1] [1] [1] [1]
H ₁₀ N ₂ O ₄ S	(NH ₄) ₂ SO ₃ • H ₂ O	ammonium sulfite hydrate	c	-283.8			[1]
H ₁₀ N ₂ O ₆ P	N ₂ H ₄ • 2H ₃ PO ₃	hydrazine diorthophosphite	c aq				
H ₁₀ N ₂ O ₆ P ₂	(NH ₄) ₂ H ₂ P ₂ O ₆	diammonium hypophosphate	c aq				
H ₁₀ N ₄ O ₄ S	(N ₂ H ₄) ₂ H ₂ SO ₄	dihydrazine sulfate	c aq m=1	-229.2 -221.0	-138.5	77	[1] [1]
H ₁₂ N ₃ O ₄ P	(NH ₄) ₃ PO ₄	tertiary ammonium orthophosphate	c aq 600 H ₂ O	-229.6 -391.3			[1] [1]
H ₁₂ N ₄ O ₅ S	(N ₂ H ₄) ₂ H ₂ SO ₄ • H ₂ O	dihydrazine sulfate hydrate	c	-291.3			[1]
H ₁₄ O ₆ S	H ₂ S • 6H ₂ O	hydrogen sulfide hexahydrate	c	-431.2			[2]
H ₁₅ O ₆ P	PH ₃ • 6H ₂ O	phosphine hexahydrate	c	-422.7			[2]
H ₁₈ N ₃ O ₇ P	(NH ₄) ₃ PO ₄ • 3H ₂ O	tertiary ammonium orthophosphate trihydrate	c	-610.8			[1]
N	N	nitrogen monatomic	g	112.979	108.883	36.622	[1]
NO	NO	nitric oxide	g	21.57	20.69	50.35	[1]
NO ₂	NO ₂	nitrogen dioxide	g	277.4			[1]
NO ₃	NO ₃	nitrogen trioxide	g	13.0			[2]
NP	PN	phosphorus nitride	g	25.043	18.453	50.437	[5]
NS	SN	sulfur nitride	g	63.000	56.278	53.055	[5]
N ₂	N ₂	nitrogen	g	0	0	45.77	[1]
N ₂ O	N ₂ O	nitrous oxide	g	19.61	24.90	52.52	[1]
N ₂ O ₃	N ₂ O ₃	dinitrogen trioxide	g	20.01 12.02	33.32	74.61	[1] [1]
N ₂ O ₄	N ₂ O ₄	dinitrogen tetroxide	g	12.19 -4.66	23.38 23.29	72.70 50.0	[1] [1]
N ₂ O ₅	N ₂ O ₅	dinitrogen pentoxide	g c	2.7 -10.3	27.5 27.2	85.0 42.6	[1] [1]
N ₅ P ₃	P ₃ N ₅	phosphorus pentanitride	c	-71.4			[1]
O	O	oxygen monatomic	g	59.555	55.388	38.467	[1]
OP	PO	phosphorus monoxide	g	-1.455	-8.391	53.219	[3]
OS	SO	sulfur monoxide	g	1.5			[1]

Table I. Selected Thermodynamic Data (Cont.)

Empirical Formula	Functional Group Formula	Name	State	ΔH_f° 298 kcal mole ⁻¹	ΔF_f° 298 kcal mole ⁻¹	S_f° 298 cal mole ⁻¹ deg ⁻¹	Ref.
O ₂	O ₂	oxygen diatomic	m=1	g aq	0 -2.8	0 3.9	48.996 26.5 [1] [1]
O ₂ P	P ₂ O	phosphorus dioxide	g	71.000	72.834	60.607	[7]
O ₂ S	SO ₂	sulfur dioxide	g ↓	-70.944 -76.6	-71.749	59.30	[1] [1]
		undissociated, m=1	aq	-77.194	-71.872	38.7	[1]
		100 H ₂ O	aq	-78.054			[1]
		200 H ₂ O	aq	-78.355			[1]
		500 H ₂ O	aq	-78.811			[1]
		1000 H ₂ O	aq	-79.201			[1]
		2000 H ₂ O	aq	-79.642			[1]
		5000 H ₂ O	aq	-80.209			[1]
		10000 H ₂ O	aq	-80.584			[1]
O ₃	O ₃	ozone	g	34.1	39.0	57.08	[1]
O ₃ S	SO ₃	sulfur trioxide	g ↓ c,β	-94.21 -105.41 -108.63	-88.69 -88.04 -88.19	61.34 22.85 12.5	[1] [1] [1]
O ₆ P ₄	P ₄ O ₆	phosphorus trioxide	c	-392.0			[1]
O ₁₀ P ₄	P ₄ O ₁₀	phosphorus pentoxide	c amorph	-713.2 -727	-644.8	54.70	[1] [1]
P	P	phosphorus, monatomic	g			38.978	[1]
		phosphorus, white, α, c III	c	0	0	9.82	[1]
		phosphorus, red triclinic	c	-4.2	-2.9	5.45	[1]
		phosphorus, black	c	-9.4			[1]
		phosphorus, red amorphous	amorph	-1.8			[1]
PS		phosphorus sulfide	g	22.500	9.694	56.033	[5]
P ₂		phosphorus, diatomic	g	34.5			[1]
P ₂ S ₃	P ₂ S ₃	phosphorus trisulfide	c	-19.2			[1]
P ₄		phosphorus tetratomic	g	14.08	5.85	66.89	[1]
P ₄ S ₃	P ₄ S ₃	phosphorus sulfide	g ↓ c	-19.408 -36.077 -37.000	-23.826 -37.513 -37.986	76.280 49.510 48.000	[3] [3] [3]
S	S	sulfur, rhombic	c	0	0	7.60	[1]
		sulfur, monoclinic	c	0.08			[1]
		sulfur monatomic	g	66.636	56.949	40.094	[1]
S ₂		sulfur diatomic	g	30.68			[1]
S ₃		sulfur triatomic	g	31.7			[1]
S ₄		sulfur tetratomic	g	32.7			[1]
S ₅		sulfur pentatomic	g	29.6			[1]
S ₆		sulfur hexatomic	g	24.5			[1]
S ₇		sulfur heptatomic	g	27.1			[1]
S ₈		sulfur octatomic	g	24.45	11.87	102.98	[1]

References

- [1] W. H. Evans, "Selected Thermochemical Values", NBS Report 8504, 1 July 1964, pp. 164-184.
- [2] F. D. Rossini, D. D. Wagman, W. H. Evans, S. Levine and I. Jaffe, "Selected Values of Chemical Thermodynamic Properties", NBS Circular 500 (U.S. Government Printing Office, Washington,, D. C. 20402 1952).
- [3] JANAF Thermochemical Data, Dow Chemical Co., Midland, Mich. June 30, 1960.
- [4] JANAF Thermochemical Data, Dow Chemical Co., Midland, Mich., Suppl. No. 1, March 31, 1961.
- [5] JANAF Thermochemical Data, Dow Chemical Co., Midland, Mich., Suppl. No. 2, June 30, 1961.
- [6] JANAF Thermochemical Data, Dow Chemical Co., Midland, Mich., Suppl. No. 6, June 30, 1962.
- [7] JANAF Thermochemical Data, Dow Chemical Co., Midland, Mich., Suppl. No. 7, September 30, 1962.
- [8] JANAF Thermochemical Data, Dow Chemical Co., Midland, Mich., Suppl. No. 8, December 31, 1962.
- [9] JANAF Thermochemical Data, Dow Chemical Co., Midland, Mich., Suppl. No. 9, March 31, 1963.
- [10] JANAF Thermochemical Data, Dow Chemical Co., Midland, Mich., Suppl. No. 13, March 31, 1964.